Fish genetic resources (FiGR) comprise all finfish and aquatic invertebrate genetic material that has actual or potential value for capture fisheries and aquaculture. In capture fisheries, more species are becoming endangered and more stocks overexploited. Management of FiGR can help maintain and rebuild these fisheries. Deep-sea fisheries and modern genetic technologies are emerging areas that require attention. Improved information is necessary for improved policies, but at present it is incomplete, scattered and unstandardized. Although tremendous progress has been made in the genetic improvement, genetic stock identification and genomics of aquatic species, further work is needed to: i) assess the status of FiGR in capture fisheries and aquaculture; ii) improve the capacities of scientists, technical persons, governments and industry; iii) improve facilities for characterizing FiGR; iv) develop genetically improved farmed types of aquatic species; v) develop appropriate policy instruments on use and conservation of FiGR; vi) improve general awareness and levels of knowledge about FiGR; and vii) prioritize species, geographic areas and production systems on which to expend resources for conservation and use of FiGR.
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Workshop on Status and Trends in Aquatic Genetic Resources

A basis for international policy

8–10 May 2006
Victoria, British Columbia, Canada

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FOOD AND AGRICULTURE ORGANIZATION OF THE UNITED NATIONS
Rome, 2007
Preparation of this document

This document represents the proceedings of the Workshop on Status and Trends in Aquatic Genetic Resources: a Basis for International Policy, held in Victoria, British Columbia, from 8 to 10 May 2006. The workshop and invited experts were supported by the FAO Commission on Genetic Resources for Food and Agriculture; organization and local support was provided by the World Fisheries Trust. The proceedings were compiled and technically edited by D.M. Bartley of the FAO Fisheries and Aquaculture Department (Rome), B.J. Harvey of the World Fisheries Trust (Canada) and R.S.V. Pullin, FAO consultant (Philippines). The cover drawing was prepared by Emanuela D’Antoni, FAO Fisheries and Aquaculture Department; general assistance was provided by Pilar González-Villegas.
Abstract

This document contains the proceedings of the Workshop on Status and Trends in Aquatic Genetic Resources: a Basis for International Policy, convened in Victoria, British Columbia, Canada, from 8 to 10 May 2006. Experts in the fields of aquaculture, biotechnology, fishery genetics, international development and international policy contributed scholarly reviews on the status of aquatic genetic resources and trends in their conservation and use in capture fisheries and aquaculture, and identified key policy issues, priorities and implications for the international development community, FAO and the FAO Commission on Genetic Resources for Food and Agriculture.

Fish genetic resources (FiGR) comprise all finfish and aquatic invertebrate genetic material that has actual or potential value for capture fisheries and aquaculture. In capture fisheries, both inland and marine, more species are becoming endangered and more stocks overexploited. Management of FiGR can help maintain and rebuild these fisheries. Deep-sea fisheries and modern genetic technologies are emerging areas that require attention. Aquaculture is expanding rapidly and now accounts for about 50 percent of the aquatic foods that are directly consumed by humans. Although genetic resources and technologies are playing a part in this expansion, they have not yet been used to extents comparable to their use in agriculture.

There is an urgent need to develop international policies for FiGR, and the breadth and complexity of capture fisheries and aquaculture present significant challenges to this process. However, the status and trends of FiGR use and conservation need to be assessed as a basis for sound policies. The workshop identified areas where further work is needed and the major activities that will be important to develop.

Information on FiGR was identified as a key issue. At present, it is incomplete, scattered and unstandardized. For wide use, information on FiGR should be global, authoritative, free and objective.

Although tremendous progress has been made in the genetic improvement, genetic stock identification and genomics of aquatic species, much further work is needed:
• to assess the status of FiGR in capture fisheries and aquaculture;
• to improve the capacities of scientists, technical persons, governments and industry;
• to improve facilities for characterizing FiGR;
• to develop genetically improved farmed types of aquatic species;
• to develop appropriate policy instruments on use and conservation of FiGR;
• to improve general awareness and levels of knowledge about FiGR; and
• to prioritize species, geographic areas and production systems on which to expend resources for conservation and use of FiGR.

Bartley, D.M.; Harvey, B.J.; Pullin, R.S.V. (eds).
Workshop on Status and Trends in Aquatic Genetic Resources: a Basis for International Policy. 8–10 May 2006, Victoria, British Columbia, Canada.
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1. Summary of the workshop

The Workshop on Status and Trends in Aquatic Genetic Resources: a Basis for International Policy, was convened in Victoria, British Columbia, Canada, from 8 to 10 May 2006 and attended by a small group of internationally recognized experts in the fields of aquaculture, biotechnology, fishery genetics, international development and international policy. The experts contributed scholarly reviews on the status of aquatic genetic resources and trends in their conservation and use in capture fisheries and aquaculture, and identified key policy issues, priorities and implications for the international development community in general and for FAO and the FAO Commission on Genetic Resources for Food and Agriculture (CGRFA) in particular.

Fish genetic resources (FiGR) comprise all finfish and aquatic invertebrate genetic material that has actual or potential value for capture fisheries and aquaculture. In capture fisheries, both inland and marine, more species are becoming endangered and more stocks overexploited. Aquaculture is expanding rapidly and now accounts for about 50 percent of the aquatic foods that are directly consumed by humans. Although genetic resources and technologies are playing a part in this expansion, they have not yet been used to extents comparable to their use in agriculture.

There is an urgent need to develop international policies for FiGR, and the breadth and complexity of capture fisheries and aquaculture present significant challenges to this process. Policies will need to address the differences between FiGR and other genetic resources, notably those for plants and livestock. These differences are due not only to the relatively recent domestication of most farmed aquatic species, but also to the large numbers of fished and farmed aquatic species and to the diversities of their aquatic environments (from the deep sea to small mountain streams) and of the production systems in which they are captured or farmed.

Policies will need to address current market forces from an increasing human population, increased environmental concerns, and improved efficiency of production and harvest. Other issues include information, management, risks and benefits, investments and awareness. Many issues here are common to both capture fisheries and aquaculture, and addressing these would benefit FiGR use and conservation in both. For example, there is a tremendous lack of information on the status and function of much of the world’s FiGR. There are also, however, significant issues that are unique to a given source of fish production; for example, the growing investment opportunities in aquaculture and the problems of governance of capture fisheries in areas beyond national jurisdiction, especially in the deep sea.

Information on FiGR was identified as a key issue. At present, it is incomplete, scattered and unstandardized. For wide use, information on FiGR should be global, authoritative, free and objective.

Although tremendous progress has been made in the genetic improvement, genetic stock identification and genomics of aquatic species, much further work is needed:
- to assess the status of FiGR in capture fisheries and aquaculture;
- to improve the capacities of scientists, technical persons, governments and industry;
- to improve facilities for characterizing FiGR;
- to develop genetically improved farmed types of aquatic species;
- to develop appropriate policy instruments on use and conservation of FiGR;
- to improve general awareness and levels of knowledge about FiGR; and
- to prioritize species, geographic areas, and production systems on which to expend resources for conservation and use of FiGR.
The workshop participants agreed that further prioritization of activities and species on which to work will be required. Nonetheless the following were judged to be of major importance:

- establishing and maintaining a directory of FiGR information sources and databases;
- compiling information on the status of FiGR for important exploited and potentially exploitable aquatic species;
- training in risk analysis with respect to FiGR conservation and use;
- identifying national and local gaps in capacity with respect to FiGR conservation and use, including special and urgent needs;
- creating Technical Guidelines for the Management of FiGR in support of the FAO Code of Conduct for Responsible Fisheries and other international instruments;
- linking existing national facilities with specific expertise in FiGR management at a regional level and creating a directory of these facilities and other service providers for conservation, characterization, genetic analysis and genetic improvement;
- reviewing existing international, regional, and national policy documents concerning FiGR;
- increasing general awareness of FiGR among the general public, resource managers and policy makers; and
- developing case studies of successful genetic improvement programmes and fisheries management that have incorporated genetic principles.
2. Background of the workshop

In 1995, the twenty-eighth session of the FAO Conference\(^1\) decided to extend the mandate of its Commission on Plant Genetic Resources to cover all components of biodiversity of relevance to food and agriculture. The result was the FAO Commission on Genetic Resources for Food and Agriculture (CGRFA), an intergovernmental body advising FAO on relevant policies and programmes. The FAO Conference recognized that approaches to plant, forestry, animal and fisheries genetic resources are different and require specialized expertise in each field, and that the implementation of the broadened mandate of the Commission should be step by step. The time has now come for the CGRFA to implement coverage of fish genetic resources (FiGR).

At its tenth session, the CGRFA agreed that its Secretariat, in cooperation with FAO’s relevant services, should submit a Multi-Year Programme of Work (MYPOW) to its eleventh session so that the Commission could implement its full mandate in the medium and longer term, including work related to fisheries. The Secretariat was asked to prepare a document on the status of the resources and needs of the various sectors, including fisheries. In response, the Fishery Resources Division (now Fisheries and Aquaculture Management Division) of the FAO Fisheries and Aquaculture Department and the CGRFA, in collaboration with the World Fisheries Trust (WFT), convened this workshop of internationally recognized experts in the fields of aquaculture, capture fisheries, molecular genetics and genomics, the deep sea, international development and aquatic conservation in order to:

- review the status of trends of aquatic genetic resources and biodiversity in capture fisheries and aquaculture (see contributed papers section); and
- identify policy issues, priorities and implications for the international development community, and specifically for FAO and the CGRFA, with regard to aquatic genetic resources and biodiversity.

3. Report of the workshop

The term fish genetic resources (FiGR) means all finfish and aquatic invertebrate genetic material that has actual or potential value for fisheries and aquaculture, including culture-based fisheries that rely on release of hatchery-bred seed to the wild. FiGR thus include DNA, genes, gametes, individual organisms, wild, farmed and research populations, species and organisms that have been genetically altered by selective breeding, hybridization, chromosome manipulation and gene transfer. The value of such genetic diversity in food production systems and in ensuring the existence and evolution of natural populations has been well established. However, policies for managing these resources at the global level are generally lacking. The report concerns almost exclusively FiGR, but farmed aquatic plant genetic resources such as seaweeds are mentioned where appropriate.

Although the CGRFA expanded its mandate to cover aquatic species in 1995, it has taken over a decade to begin to address relevant issues. Workshop participants expressed a sense of urgency for the development of adequate policies for the sustainable use and conservation of FiGR. In both inland and marine capture fisheries, more species are becoming endangered and more stocks over-exploited. Currently, about 50 percent of the aquatic foods consumed by humans come from aquaculture.

FiGR are valuable not only because of their importance in aquaculture and the need to accelerate genetic improvement of farmed aquatic populations, but also because wild stocks are under threat and declining, and wild gene pools represent and ensure the continued survival of populations and species.

Although there are international and regional institutions and organizations that are contributing to addressing these problems (Table 1), there is no global strategy for the management — i.e. the conservation and use — of FiGR. Specific strategies are required for in situ conservation of FiGR on farms and in natural ecosystems, and for ex situ conservation of FiGR and as cryopreserved gametes or embryos.

3.1 SPECIAL CHARACTERISTICS OF FiGR AND THE AQUATIC ENVIRONMENT

In 1995, the twenty-eighth FAO Conference recognized that different approaches are needed for managing plant, forestry, animal and fisheries genetic resources. The domestication of most of the aquatic species used in aquaculture has a much shorter history than the domestication of plant and livestock species in agriculture and there are many other unique features of the aquaculture and fisheries sectors with respect to conservation and use of genetic resources. The workshop identified the following special features of aquatic species and FiGR that should be considered in policy development:

- Most species of farmed fish have a relatively short history of domestication and genetic improvement.
- Some species of farmed fish have reproductive characteristics (very high fecundity and short generation times) that can facilitate rapid genetic improvement.

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2 Pullin et al., 1999.
3 Bartley and Toledo, this volume; Pullin, this volume.
4 Grant, this volume; Smith, this volume; FAO, 2004, http://www.fao.org/DOCREP/007/y5600e/y5600e00.htm
6 Pullin, this volume.
Some international and regional initiatives that address aquatic genetic resources in capture fisheries and aquaculture

<table>
<thead>
<tr>
<th>Activity</th>
<th>Theme</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>1985 International Association for Genetics in Aquaculture</td>
<td>Genetic improvement in aquaculture</td>
<td>Numerous peer-reviewed publications and network of geneticists</td>
</tr>
<tr>
<td>1992 United Nations Conference on Environment and Development</td>
<td>Sustainable use and conservation of FiGR, plus fair and equitable sharing of benefits</td>
<td>Legally binding International Convention on Biological Diversity; Agenda 21; Jakarta Mandate on Marine and Coastal Biodiversity; Cartagena Biosafety protocols</td>
</tr>
<tr>
<td>Development of Genetically Improved Farmed Tilapia (GIFT)</td>
<td>Use of traditional animal breeding and selection from a diverse gene pool to create faster growing and hardier fish for developing-country aquaculture</td>
<td>Successful, public-funded projects that produced GIFT and disseminated them in Asia and the Pacific to become the basis of national tilapia breeding programmes; the projects also developed capacity for national breeding programmes in Asia and Africa</td>
</tr>
<tr>
<td>1993 International Network for Genetics in Aquaculture</td>
<td>Enhancing research and developing collaborative linkages that could help establish national breeding programs</td>
<td>Facilitated development of fish breeding programmes, exchange of information, assessment of genetic improvement programmes and capacity building</td>
</tr>
<tr>
<td>1995 FAO Code of Conduct for Responsible Fisheries</td>
<td>Sustainable fisheries and aquaculture in an environmentally and socially acceptable manner</td>
<td>Soft law code used by FAO member States et al. for fisheries development and management; Technical Guidelines in support of the Code have been produced for many areas of capture fisheries and aquaculture - none yet for FiGR</td>
</tr>
<tr>
<td>1995 FAO Commission on Genetic Resources for Food and Agriculture</td>
<td>Inter-governmental body to formulate policies on genetic resources in, inter alia, capture fisheries and aquaculture</td>
<td>The Commission has not yet addressed FiGR</td>
</tr>
<tr>
<td>2001 FAO Committee on Fisheries Sub-Committee on Aquaculture</td>
<td>Inter-governmental forum to address all issues relevant to aquaculture development and management; its parent body is the Committee on Fisheries</td>
<td>Provides advice to FAO Fisheries and Aquaculture Department, but has addressed FiGR so far only in general terms of sustainable use</td>
</tr>
<tr>
<td>2004 ICES Code of Practice on the Introductions and Transfers of Marine Organisms</td>
<td>Code on procedures and risk assessment for the introduction of alien species that also includes genetically altered organisms</td>
<td>Code of practice that has been adopted in principle by FAO and FAO Regional Bodies</td>
</tr>
</tbody>
</table>

1 [http://www.medialaqua.fr/AGA/webgeneral_information/index.htm](http://www.medialaqua.fr/AGA/webgeneral_information/index.htm)
4 [http://www.worldfishcenter.org/nga/network.htm](http://www.worldfishcenter.org/nga/network.htm)

The variety of aquatic species that are fished and farmed is very high.
Fished and farmed aquatic species have very diverse life histories including, for example, short- and long-lived species.
The wild relatives of farmed aquatic species are also important for future breeding programs in aquaculture.
Some farmed aquatic species that escape from captivity can readily establish feral populations.

The workshop also noted that aquatic production systems, species and environments have the following special features:
- Production systems include not only conventional capture fisheries that target wild stocks and aquaculture that is based on farming captive-bred fish, but
also culture-based fisheries that are stocked from hatcheries and capture-based aquaculture in which wild-caught fish are fattened.

- Some aquatic species that are fished or farmed are used in recreational and ornamental fisheries.
- Some aquatic species and their wild populations are seriously threatened with genetic change or extinction.
- Distinct types of farmed aquatic species are generally less threatened, but become so as farmers choose to retain only the most recently developed and profitable.
- Some threatened and endangered fish species are being targeted by capture fisheries or taken as bycatch.
- The numbers of farmed aquatic breeds/strains/varieties and other types are increasing.
- Almost all aquatic species that are hunted and trapped in capture fisheries are wildlife, and are often regarded as common property resources.
- Capture fisheries may take place in open access environments or in areas not under national or international jurisdiction, e.g. high seas.
- Capture fisheries and aquaculture often impact and are themselves impacted by other users of natural resources, especially inland waters (irrigated agriculture/domestic and industrial use), forestry, human settlements, tourism, and waste disposal.
- **Ex situ** and **in situ** conservation of FiGR are important, but can be difficult and costly.
- Aquatic environments in capture fisheries are extremely diverse, from the deep sea to mountain streams, and are also typically difficult to monitor.
- Aquatic environments in aquaculture range from highly controlled intensive recirculation systems to open water cage, pen, pond and raceway systems in fresh, brackish and marine waters and in most temperature zones.
- Aquatic environments in capture fisheries and aquaculture are often interconnected. In particular some capture fisheries take place in waters that are transboundary, international, and sometimes beyond the scope of any effective jurisdiction.

### 3.2 DRIVERS INFLUENCING MANAGEMENT OF FiGR

In order to develop appropriate policies on FiGR, key drivers influencing their management need to be identified. “Drivers” refers to trends that influence the conservation and sustainable use of FiGR. The workshop identified the following key drivers.

**Driver 1: Market forces**

- increased demand for food fish due to human population growth, increased affluence and the many health benefits of fish will increase pressure on farmed and wild populations;
- globalization and competition for markets within and among food production sectors will stimulate competition for aquatic resources and necessitate good marketing;
- competition for inputs, resources and space will force fish production to be more cost-effective and efficient; and
- consumer attitudes to some aquatic food production systems and to some new technologies (for example, genetically improved farmed fish and farming systems that are perceived as environmentally and/or ethically unsound) will constrain their adoption.
Driver 2: Environmental issues
- stagnation and decline of capture fisheries due to overexploitation and habitat degradation will force improved management in some cases and increase reliance on aquaculture or alternative foods in others;
- increased environmental awareness on the part of policy makers and the public will result in increased demand for sustainable use of fishery products;
- availability of fresh water will change in response to climate change and the needs of human population growth and development; and
- climate change will alter the potentials for capture fisheries and aquaculture in some areas and FiGR are the basis for sustaining the ability of aquatic species to adapt to changed environments, in nature and in farming systems.

Driver 3: Production and management forces
- alien aquatic species and genotypes will present opportunities (increased production and value) and problems (loss of wild biodiversity and habitat);
- because most capture fishery resources have been fully explored and there are few new species or areas available, better management of existing stocks or increased reliance on other food sources will be required;
- improved methods for fishing and farming will enable the sectors to expand;
- issues of sustainability have arisen in both capture fisheries and aquaculture and improved methods of fishing and farming are needed to sustain or expand production;
- scientific advances, particularly in the application of genetic technologies, including genomics, to capture fisheries and aquaculture, will provide opportunities for improved fish production;
- intensification of farmed fish production and harvest systems will produce more food per unit area and require improved breeds and management;
- access to FiGR, benefit sharing and intellectual property rights will influence use and policies; and
- increasing consolidation of farmed fish production systems with feed and seed suppliers is likely to have different effects on large- and small-scale producers.

3.3 ISSUES INFLUENCING MANAGEMENT OF FiGR
The breadth and complexity of the fishery and aquaculture sectors present significant challenges to the development of international policies on FiGR. Addressing the wide range of issues and special features of FiGR will take time and substantial human and financial resources. The mandate of this workshop was to present an unprioritized range of issues to the CGRFA. Prioritization of species on which to work, geographic areas, and production systems etc. Will be the work of future fora convened to develop specific details of the MYPOW or other programmes of work.

The issues presented below concern information, management, risks and benefits, investments, awareness, and policy. Some FiGR issues here are common to capture fisheries and aquaculture; for example, some wild FiGR of importance for both capture fisheries and aquaculture are being overfished. There are also important FiGR issues that are specific to either capture fisheries or aquaculture; for example, the difficulties of capture fisheries governance in high seas and areas beyond national jurisdiction, and the growing investment opportunities in aquaculture.

Issue 1: Information (see also section 3.4)
For both capture fisheries and aquaculture, there are gaps in information on the status of FiGR and on trends in their conservation and use. Information is often scattered, incomplete and not easily accessible. Genetic information about fish populations is often limited. Where population genetic data do not exist or are too expensive or difficult to
collect, especially in some developing countries, surrogate criteria and indicators can sometimes be developed to predict genetic stock structure or to identify genetically unique populations or strains. For example, within a given species, populations that exhibit different life histories, have different migration times, or inhabit different river basins can be expected to be genetically different.

In capture fisheries, lack of information about fish stocks leads to a lack of regulation and to illegal, unreported and unregulated (IUU) fishing. Information is increasing for a change to ecosystem-based management of capture fisheries, but the importance of FiGR and other genetic resources in ecosystem function are yet not well understood. Most important, existing genetic information on fish stocks is often simply not used in fishery management.

Issue 2: Management of FiGR
Capture fisheries and aquaculture share several FiGR management issues. Because of a lack of consensus on global priorities, fisheries development and conservation programmes remain largely divorced from FiGR management concerns. Ownership of and access to FiGR, and sharing the costs of FiGR conservation and the benefits from FiGR use, are also issues for both capture fisheries and aquaculture.

Management – i.e., conservation and sustainable use – of FiGR is often ignored in capture fisheries. This applies not only to the target species but also to key species for ecosystem function and to bycatch species, which are often more vulnerable to extinction than the target species. Capture fisheries can damage habitats, thereby endangering biodiversity, including marine mammals and seabirds. Capture fisheries can have particularly severe impacts on populations of slow growing or late maturing species.

In aquaculture, objectives of development or of assistance are often not clearly defined, resulting in confusion between farming for local food security and farming for export. The wild relatives of farmed fish have actual or potential value, and are often important as food sources in developing countries, so their stewardship must be adequately compensated. There are at present few international efforts to conserve the wild relatives of farmed aquatic species.

Issue 3: Genetic risks and benefits
Capture fisheries confront basic conceptual problems such as the definitions of “population” and “stock” – key concepts in the analysis of genetic risk. In aquaculture, there is a need for cost/benefit analysis of breeding programmes and genetic resources management. The use of alien species and alien genotypes in aquaculture and stocking programmes is unevenly regulated in developed and developing countries alike, and the consequent risks to wild and farmed populations are not quantified. Movement of stocks, introductions and transfers, and interactions between hatchery and wild stocks as a result of escapes or deliberate release have yet to be well analysed in terms of their risks to wild and farmed FiGR. To deal with biosafety issues, genetic risk assessment based on genetic stock identification, especially for culture-based fisheries and capture-based aquaculture, was identified as a high priority. Guidelines or codes of conduct on genetic resource management would be useful in addressing many of the management and risk/benefit concerns.

Issue 4: Investments and applications
FiGR conservation and use in aquaculture presents significant investment opportunities. However, genetic improvement strategies in aquaculture, from domestication and selective breeding to hybridization and other forms of genetic alteration, can be applied only where there are adequate resources, in terms of human and institutional capacities and prioritized funding. As aquaculture produces more of world’s fish supply, the
value of FiGR for farmed and potentially farmable fish is increasing, but this has not yet been recognized in terms of increased investment in their management.

**Issue 5: Education and awareness**
In capture fisheries and aquaculture, decision makers often fail to appreciate the urgency to act before species or valuable stocks/strains go extinct. There is also widespread consumer ignorance of how food fish are produced, and most of the general public have no concept of FiGR.

Many capture fisheries professionals are also unaware of the importance of FiGR. In developed and developing counties, many fisheries policymakers and managers either do not know how to use genetic information when it does exist, or are unaware of its existence.

In aquaculture, professional awareness of the importance of FiGR is relatively high in the developed world and increasing in developing countries, but everywhere there is little public awareness about how farmed fish are bred and sometimes misinformation about the actual and potential applications of genetics in aquaculture (Liu, 2007).

**Issue 6: Policy instruments and mechanisms**
While policies on FiGR are lacking or inadequate for most capture fisheries, the problem is especially acute with deep sea fisheries. In aquaculture, advances in molecular biology and genetics are outpacing policy formulation for their application and regulation. Policies regulating use of FiGR and alien species/genotypes, when they exist, are often difficult to enforce. The genetic resources of farmed aquatic plants are a special case, as they are not yet adequately covered by existing instruments for plant genetic resources or as FiGR.

Capture fisheries and aquaculture in general lack adequate FiGR policy instruments, at international, regional, national and local levels. This reflects the ongoing inadequacies of efforts to document and to monitor FiGR and to provide for the sharing of costs for their conservation and of benefits from their use, especially for poor people. In developing countries, inadequate human capacity and infrastructure, including low capacity for risk assessment and management when using genetically altered forms, are especially acute. In general, policy formulation will need to balance a cross-sectoral, multidisciplinary approach (that addresses poverty alleviation and FiGR conservation) with more focussed approach to address specific topics, such as genetic improvement in aquaculture.

### 3.4 FiGR INFORMATION SOURCES AND NEEDS

FiGR information refers broadly to genetic characterization (e.g. genetic sequences and other measures of genetic diversity at individual and group levels), breeding histories, performance data, and behavioural and life cycle characteristics. Categories of FiGR information include: DNA; genes; gametes; individual organisms; wild, farmed and research populations; species; forms that have been genetically altered by selective breeding, hybridization, chromosome manipulation and gene transfer; and methods for genetic characterization, FiGR conservation, and genetic improvement.

For wide use by the Members of FAO and others, FiGR information should be global, authoritative, free and objective. At present, much FiGR information is incomplete, scattered and held in diverse formats. No existing databases give adequate coverage to FiGR or consolidate existing information, although there are some excellent information sources for specific topics; for example, FishBase\(^7\) has good coverage of cytogenetics and some population genetics. The National Institutes of

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\(^7\) [www.fishbase.org](http://www.fishbase.org)
Health of the United States of America maintains genomic databases on molecular genetics and bioinformatics.\(^8\)

Current FAO datasets on capture fisheries and aquaculture, include very little information on FiGR. The FAO Species Fact Sheets on farmed aquatic species contain good information on taxonomic features and natural history, but coverage on their genetics is uneven and often lacking. As the number of farmed fish strains, hybrids, and other genetically altered forms increases in aquaculture, aquaculture statistics will need to capture their relative contributions to farmed fish production and value, as is done for livestock.\(^8\) This would assist both conservation and use of FiGR. Similarly, fuller information on the genetics of wild fish populations would improve their conservation and use as FiGR for capture fisheries and aquaculture.

In order to initiate and develop its coverage of FiGR, the CGRFA can draw upon its long experience with plant, and to a lesser extent livestock genetic resources information that is of importance to FAO Member States for policy-making and management. FiGR information is held by diverse groups in the public and private sectors. The CGRFA will have to consider to what extent it might need to become itself a centre for FiGR information that FAO will collect and hold, as well as offering linkages with and portals into FiGR information sources collected and held by others. The latter, decentralized system already exists to a limited extent, but much existing FiGR information has limited accessibility because of non-standardized formats and terminology and its reliability and provenance are rarely well checked.

FAO fish production statistics, from capture fisheries and aquaculture, are standardized and represent official government information, but have almost no information regarding FiGR. For CGRFA coverage of FiGR, the use of other sources of FiGR information that are not the official reports of its Members should not be a problem, provided that information meets the criteria of authoritativeness and objectivity stated above.

The workshop appreciated that gathering, compiling and disseminating information on FiGR will require human and financial resources. Therefore, it will be necessary to convince the collectors and holders of FiGR information such as international, regional, national and local organizations, that their FiGR information is useful and that making it more widely available as part of FAO’s global coverage of FiGR will be of mutual benefit. Provision of FiGR information and facilitating linkages to FiGR information sources will help the Members and partners of FAO to:

- fulfil obligations under international conventions such as Convention on Biological Diversity, FAO Code of Conduct for Responsible Fisheries (see previous footnotes), and the Convention on International Trade in Endangered Species of Fauna and Flora (CITES);\(^10\)
- facilitate better management of their FiGR through shared information and experiences;
- improve the identification and traceability of aquatic produce;
- assist risk assessment associated with the movement of aquatic species and the use of genetically altered species;\(^11\)
- secure funding and cooperation from donors and partners; and
- seek compensation for adverse impacts on FiGR.

\(^8\) http://discover.nci.nih.gov/
\(^10\) http://www.cites.org/
\(^11\) Genetic alteration may be the result of a number of genetic technologies, including hybridization, selective breeding, chromosome set manipulation, genetic engineering and gene transfer.
4. Conclusions and recommendations of the workshop

Tremendous progress has been made in the fields of fish genetic improvement (Liu, 2007; Pullin, 2007), genetic stock identification (Grant, 2007; Smith, 2007) and genomics (Liu, 2007). The stage is clearly set for the creation of policies on FiGR that reflect this body of experience and anticipate future global needs, especially in view of the expansion of aquaculture and the decline in many wild aquatic populations. The FAO Fisheries and Aquaculture Department, CGRFA and partners will be expected to play major roles in this area over the next several years.

The material presented in this summary and in the following review papers represents scientific analyses of extremely diverse, complex and sometimes controversial topics. Policies for the management of the world’s FiGR will depend on a variety of factors. Work plans of the CGRFA will need to reflect that variety. It is the workshop participants’ hope and recommendation that other fora, including those organized by the CGRFA, will find this material useful for prioritizing areas for future work, in order to meet global development and conservation objectives. Prioritization will need to consider, inter alia, species, production systems, geographic coverage, risks and benefits associated with different technologies, consumer perspectives and ethics.

Pending this prioritization, the workshop participants recommended the following next steps toward developing policy instruments on the use and conservation of FiGR:

- assess the status of FiGR in fisheries and aquaculture;
- identify and fill regional capacity needs for scientists, technical persons, government and industry;
- improve facilities for characterizing FiGR;
- continue genetic improvement of farmed aquatic species;
- improve general awareness and knowledge of FiGR;
- assess existing FiGR policy instruments; and
- explore the twinning (i.e. co-planning, co-financing, co-governance) of aquaculture operations with conservation of wild aquatic genetic resources and related habitats.

These recommendations are elaborated upon below.

4.1 ASSESS THE STATUS OF FiGR

FiGR exist “in situ and in vivo” (as free-living, wild and feral populations, and as captive populations on-farm), “ex situ and in vitro” (as collections of cryopreserved sperm, embryos and other tissues/DNA), and “ex situ and in vivo” (as aquarium and research populations). Increasing the amount and quality of information on the status of FiGR could use updatable geographic information systems that incorporate genetic information, including diversity and abundance measures. A consultation on existing databases could be convened in order to assess their ability to incorporate this extraordinary diversity. Several good general information sources exist (Pullin, 2007; Liu, 2007; Smith, 2007), as well as specialized databases on key species, e.g. common carp, or groups of species such as Pacific salmon and tilapia. A directory of information sources and databases is needed, and establishing and maintaining such a directory could be suitable roles for the CGFRA and the FAO Fisheries and Aquaculture Department.
The status of important farmed aquatic species groups (including tilapias, carps, catfishes, penaeid shrimps, bivalves, abalones, seaweeds, and freshwater macrophytes) could be compiled, reviewed and synthesised. For marine capture fisheries, the most important groups include small pelagics, reef fishes, elasmobranches, large pelagics, demersals, and diadromous fishes. Important inland capture fisheries groups include those for many of the farmed species (such as carps, catfishes, characins, cichlids and salmonids), as well as many others described under the International Statistical Standard Classification of Aquatic Animals and Plants (ISSCAAP) scheme. With such a large array of species to study, clear prioritization and working through partnerships will be necessary. Documentation of the status of FiGR for these groups can link to other information sources such as FishBase, the FAO cultured species fact sheets and the FAO Species Identification Programme. Work has already begun on summarizing the information available on salmon and trout genetic resources.

4.2 IDENTIFY AND FILL REGIONAL CAPACITY NEEDS
Capacity building should be increased to include FiGR characterization and management, breed improvement, analysis of genetic data and training in risk analysis. Well-trained persons are already engaged in characterizing FiGR in fisheries and aquaculture (see, for example, the publications of the International Association on Genetics in Aquaculture (IAGA) but they and their organizations merit more support to expand training activities. For example, training in risk analysis techniques would help those developing fish breeding programmes to make good choices of broodstock and genetic improvement techniques to meet their objectives surely and safely.

Regional networks can also play an important role in building and maintaining capacity and communication, e.g. Network of Aquaculture Centres in Asia and the Pacific (NACA), and the International Network for Genetics in Aquaculture (INGA). The Southern African Botanical Diversity Network, funded by the Global Environmental Facility (GEF) to improve information and capacity on plants, could be a useful model for regions and organizations requesting support for FiGR. The Network of Aquaculture Centres in Eastern Europe (NACEE) has recently been set up with support from FAO and could be expected to help address capacity building on FiGR, especially as capacity to improve and manage FiGR in carp and other freshwater species of commercial importance is well advanced in several member countries. Gaps in capacity should be examined on a geographic scale to identify any special regional and national needs.

FAO could consider creating Technical Guidelines for the Management of FiGR in support of the CCRF. Semi-technical manuals and scientific publications reviewing basic methods of breed improvement and methods of characterization and management of natural fish populations already exist and could be useful models.

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12 http://www.fishbase.org
15 Harvey, Brian in press FAO website.
16 www.mediaqua.fr/IAGA/web/general_information/index.htm
17 www.enaca.org
18 http://www.agrowebcee.net/subnetwork/nacee/
4.3 IMPROVE FACILITIES FOR CHARACTERIZING FiGR

New facilities in support of use and conservation of FiGR will not be necessary in all countries. Economies of scale are such that numerous small facilities analyzing small amounts of genetic material may not be economically justifiable. Improvements in transportation and communication are making collaboration among organizations cheaper and easier. Some existing facilities, together with the expertise of their staff, could be linked at the regional level. A directory of service providers for breed improvement, genetic characterization and genetic conservation could be created to facilitate access to expertise and technology and to prevent unnecessary duplication of efforts.

4.4 IMPROVE AWARENESS OF FiGR

Awareness of the importance of FiGR remains extremely poor and extends from the general public, to resource managers and through to policy makers. This is not altogether surprising, given the rapid developing state of development of genetics and its poor coverage in some school curricula, but it must be remedied as soon as possible. The first steps are to compile a list of target audiences that need specific information, then to identify appropriate channels and formats.

Part of the problem is the inability of many geneticists to communicate clearly about FiGR to the public and to professionals who are not geneticists. It was suggested that a workshop be convened to identify target audiences for learning about FiGR and to explore how best to reach them. This workshop could include participants from FAO, donors, government resource officers, Non-Governmental Organizations (NGOs), and other development groups. The International Development Research Centre of Canada has agreed to provide funding for such a workshop.

FAO should consider including an article to increase awareness of the value of FiGR and specifically to discuss the necessity of reporting on breeds/strains/stocks/hybrids in the 2008 edition of the FAO flagship publication, *State of World Fisheries and Aquaculture* (SOFIA). If information on genetic resources is to be provided to FAO, then the FAO Fisheries and Aquaculture Department, with assistance from partners, will need to provide some standardization and guidance on appropriate terminology and reporting. The reviews listed under Status above can also be included in SOFIA and used to improve awareness of policy makers, various commissions, fishery managers, hatchery managers, farm managers, industry associations, NGOs, researchers and teachers.

Case studies were proposed as a way of demonstrating the value of FiGR in fisheries and aquaculture. The Network of Aquaculture Centres in Eastern Europe (see footnote 18), the long-standing work on genetic improvement of common carp at the Fish Culture Research Institute in Szarvas, Hungary, and the well-established development, use, dissemination and management of common carp genetic resources in eastern Europe were suggested mechanisms and material for a case study. The history of the development and impact of the Genetic Improvement of Farmed Tilapia (GIFT) was also suggested. Compilation of those fisheries that are managed at the genetic stock or strain level, and those farms or areas that report production by breed, would be useful in order to better understand the practicalities, costs and benefits of collecting information on FiGR.

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20 The International Development Research Centre of Canada in collaboration with the World Fisheries Trust (Canada) subsequently convened a workshop, *Sink or Swim: Roundtable on Aquatic Genetic Resources*, Victoria, B.C. September 26/27, 2006. www.worldfish.org

21 The United Nations General Assembly recently made a similar request that FAO should look a means to revise marine capture fishery statistics based on stock structure http://daccessdds.un.org/doc/UNDOC/GEN/N04/477/70/PDF/N0447770.pdf?OpenElement

22 http://www.worldfishcenter.org/reshigh01_3.htm
4.5 ASSESS EXISTING POLICY INSTRUMENTS
Although FiGR are not well covered by most existing international, regional and national policies, any relevant policies that do exist should be appraised for their application to FiGR. Specific documents recommended for review were the FAO Code of Conduct for Responsible Fisheries (FAO, 1995), the Cartagena Biosafety Protocols and their parent, the Convention on Biological Diversity (Secretariat CBD, 2000). General documents on ownership, access and intellectual property rights should also be reviewed, especially the material transfer agreements and germplasm acquisition agreements currently used by INGA, the Consultative Group on International Agriculture Research (CGIAR), and others. Policy formulation will need to balance an holistic approach involving cross-sectoral and multidisciplinary policies on such issues as economic development, poverty alleviation and land use, with more specialized policies on FiGR that would address primarily fisheries and aquaculture, for example, public-private partnerships. The Convention on Biological Diversity develops work plans for types of ecosystems, e.g. inland waters, mountains, and deserts, whereas much of the work of FAO and the CGIAR centres is focused on geographic areas, climatic zones and specific commodity groups. The CGIAR centre with responsibility for capture fisheries and aquaculture is the WorldFish Center; the CGIAR’s Bioversity International acts as a Member-Coordinator for a System-Wide Genetic Resources Programme, which includes some coverage of FiGR.

4.6 EXPLORE THE TWINNING OF AQUACULTURE OPERATIONS AND CONSERVATION
Aquaculture operations have usually had adversarial relationships with other uses of natural resources, especially nature conservation. This is to some extent unavoidable and it applies also in much of agriculture, forestry and other development. With aquaculture now in a rapid phase of growth, particularly in the developing world where most FiGR are also located, the time is ripe to explore to what extents aquaculture operations can be planned and conducted in harmony with nature conservation, including conservation of FiGR. Reconciliation between the needs of aquaculture operations and the needs of nature conservation is sorely needed. One approach could be to twin indefinitely the financing and conduct of aquaculture operations with those of nature conservation. This would mean setting aside conservation areas that are off-limits to aquaculture and to all contact with farmed fish and farm waters. Some potential sites for this already exist as nature reserves, sacred groves, etc. Similarly the practice of establishing aquatic protected areas is becoming a key part of capture fisheries management in many areas. For aquaculture production, the pay-offs would be not only the survival of threatened wild FiGR of present or likely future importance for breeding programmes, but also a platform from which to argue for permission to use, in designated farming areas, the most profitable species and genetically altered farm types available — as is the case for most of agriculture.

24 http://www.biodiv.org/biosafety/default.aspx
25 http://www.worldfishcenter.org/inga/
26 see for example http://www.ciat.cgiar.org/improved_germplasm/mta_breeding.htm
27 www.worldfishcenter.org
28 www.bioversityinternational.org
Annex 1

Agenda

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<th>Monday 8 May</th>
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<td><strong>08.30 – 11.00</strong></td>
<td>Session I: Welcome and objectives of workshop</td>
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<td>08.30</td>
<td>Registration</td>
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<td>09.00 – 10.00</td>
<td>Welcome by World Fisheries Trust</td>
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<td>Welcome by FAO</td>
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<td>Introduction to workshop agenda &amp; objectives</td>
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<td>10.00 – 10.30</td>
<td>Overview of FAO Fisheries, the Code of Conduct for Responsible Fisheries and the CGRFA*</td>
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<td>Capture Fisheries</td>
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<tr>
<td><strong>09.00 – 15.00</strong></td>
<td>Session III: Synthesis of reviews</td>
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<td>Genomics and Modern Genetic Technologies</td>
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<td>John Liu</td>
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<td>09.00 – 12.00</td>
<td>Identification of key issues: technical and policy</td>
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<td>Facilitated discussion and drafting</td>
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<td>12.30 – 14.00</td>
<td>Lunch</td>
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<td>Coffee</td>
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<td>15.30 – 17.00</td>
<td>Session IV: Priorities for action</td>
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<td>Identification of strategic actions in both policy and technical areas</td>
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<td>Facilitated discussion and drafting</td>
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<th>Wednesday 10 May</th>
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<td><strong>09.00 – 17.00</strong></td>
<td>Session V: Finalize report</td>
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Annex 2

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CONTRIBUTED PAPERS
Developing policies for the management of fishery genetic resources

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1. SUMMARY
Policy on aquatic genetic resources is primarily guided in FAO by the Code of Conduct for Responsible Fisheries and the Convention on Biological Diversity. These instruments are complementary and both recognize the importance of sustainable use and conservation of aquatic genetic resources. Policies will be influenced by a number of drivers that include the increasing human population, resource limitations, the need to address broad and complex social issues, intensification of farming and fishing systems, increases in technology, and the recognition of sovereign rights of countries in regards to aquatic genetic resources.

2. INTRODUCTION
The Preamble to the 1989 edition of the Constitution of the Food and Agriculture Organization of the United Nations defines the common purpose of the Nations accepting the Constitution as:
- raising levels of nutrition and standards of living of the peoples under their respective jurisdictions;
- securing improvements in the efficiency of the production and distribution of all food and agricultural products;
- bettering the conditions of rural populations; and thus
- contributing toward an expanding world economy and ensuring humanity’s freedom from hunger.

The Fisheries and Aquaculture Department of FAO promotes sustainable and responsible fisheries through its work to improve policy, legislative and institutional frameworks, to develop and evaluate technologies in fisheries and aquaculture, to build capacity and to collect and disseminate information on the world’s fisheries and aquaculture. In 1995 the FAO Council adopted the FAO Code of Conduct for Responsible Fisheries (CCRF) (FAO, 1995) that has since become the framework and primary mechanism through which Member Governments have addressed the above issues. The vision of the Fisheries and Aquaculture Department is: A world in which responsible and sustainable use of fisheries and aquaculture resources make an appreciable contribution to human well-being, food security and poverty alleviation. Working through Governments and appropriate Ministries, the Fisheries and Aquaculture Department acknowledges a focus on fishers and fish farmers.
Collecting information on the status and trends of aquatic genetic diversity is extremely difficult, especially for global repositories of this information such as FAO. The FAO Fisheries and Aquaculture Department receives yearly information on fisheries and aquaculture production from Member Countries. Although this data set represents the best available scientific information it is far from complete and includes virtually no information below the species level. Indeed much of the reported information is not identified to species (especially true for inland fishery resources). A disturbing trend is that the quantity of production not reported at the species level is increasing (FAO, 2004). Countries are better at reporting aquaculture production by species, but not by strain, breed, or variety. Thus, we have scant global information on the numerous breeds of carp, catfish, tilapia and other genetically altered species that comprise aquaculture production.

Management of the resources and collection of information from areas beyond national boundaries are further complicated by problems of governance and jurisdiction. Regional fishery bodies have been established in some marine and inland areas. However, there are gaps in coverage and problems with implementation of regional agreements.

3. CODE OF CONDUCT FOR RESPONSIBLE FISHERIES (CCRF) AND OTHER INTERNATIONAL MECHANISMS
The CCRF is a voluntary, non-binding international instrument that the Members of FAO have pledged to help implement as appropriate and to the best of their abilities. Articles of the CCRF relevant to FiGR include:

- **Article 6.2** – Fisheries management should promote the maintenance of the quality, diversity and availability of fishery resources in sufficient quantities for present and future generations in the context of food security, poverty alleviation and sustainable development. Management measures should not only ensure the conservation target species but also of species belonging to the same ecosystem or associated with or dependent upon the target species.

- **Article 7.2.2** – ...biodiversity of aquatic habitats and ecosystems is conserved and endangered species are protected.

- **Article 9.1.2** – States should promote responsible development and management of aquaculture, including an advance evaluation of the effects of aquaculture development on genetic diversity and ecosystem integrity, based on best available scientific information.

- **Article 9.3.1** – States should conserve genetic diversity and maintain integrity of aquatic communities and ecosystems by appropriate management (in particular to minimize adverse impacts from non-native and genetically altered species).

- **Article 9.3.3.** – States should ...encourage the adoption of appropriate practices in the genetic improvement of broodstock, ....

- **Article 9.3.5** – States should, where appropriate, promote research and, when feasible, the development of culture techniques for endangered species to protect, rehabilitate and enhance their stocks, taking into account the critical need to conserve genetic diversity of endangered species.

- **Article 12.8** – States should conduct research into, and monitor, human food supplies from aquatic sources ...and ensure that there is no adverse impact on consumers.

The Fisheries and Aquaculture Department works in close association with a variety of international mechanisms and agencies. The key mechanism relevant to the issue of aquatic genetic resources and biodiversity is the Convention on Biological Diversity (CBD). The FAO CCRF, as well as the CGRFA, have similar principles with, and are complementary to the CBD. Key sections of the CBD that pertain to aquatic genetic resources and biodiversity are:

- **Article 6** – Each Contracting Party shall, in accordance with its particular conditions and capabilities: (a) Develop national strategies, plans or programmes for the conservation and sustainable use of biological diversity or adapt for this
Developing policies for the management of fishery genetic resources

purpose existing strategies, plans or programmes which shall reflect, inter alia, the measures set out in this Convention relevant to the Contracting Party concerned; and (b) Integrate, as far as possible and as appropriate, the conservation and sustainable use of biological diversity into relevant sectoral or cross-sectoral plans, programmes and policies.

• Article 7 – Monitoring: (a) Identify components of biological diversity important for its conservation and sustainable use having regard to the indicative list of categories set down in Annex I; (b) Monitor, through sampling and other techniques, the components of biological diversity identified pursuant to subparagraph (a) above, paying particular attention to those requiring urgent conservation measures and those which offer the greatest potential for sustainable use; (c) Identify processes and categories of activities which have or are likely to have significant adverse impacts on the conservation and sustainable use of biological diversity, and monitor their effects through sampling and other techniques; and (d) Maintain and organize, by any mechanism, data, derived from identification and monitoring activities pursuant to subparagraphs (a), (b) and (c) above.

• Article 8 – In situ conservation: (g) Establish or maintain means to regulate, manage or control the risks associated with the use and release of living modified organisms resulting from biotechnology which are likely to have adverse environmental impacts that could affect the conservation and sustainable use of biological diversity, taking also into account the risks to human health; (h) Prevent the introduction of, control or eradicate those alien species which threaten ecosystems, habitats or species; (i) Endeavour to provide the conditions needed for compatibility between present uses and the conservation of biological diversity and the sustainable use of its components.

• Article 9 – Ex situ conservation: (a) Adopt measures for the ex-situ conservation of components of biological diversity, preferably in the country of origin of such components; (b) Establish and maintain facilities for ex-situ conservation of and research on plants, animals and micro-organisms, preferably in the country of origin of genetic resources; (c) Adopt measures for the recovery and rehabilitation of threatened species and for their reintroduction into their natural habitats under appropriate conditions; (d) Regulate and manage collection of biological resources from natural habitats for ex-situ conservation purposes so as not to threaten ecosystems and in-situ populations of species, except where special temporary ex-situ measures are required under subparagraph (c) above …

• Article 10 – Sustainable use: (b) Adopt measures relating to the use of biological resources to avoid or minimize adverse impacts on biological diversity; (c) Protect and encourage customary use of biological resources in accordance with traditional cultural practices that are compatible with conservation or sustainable use requirements; (d) Support local populations to develop and implement remedial action in degraded areas where biological diversity has been reduced.

• Article 15 – Access to genetic resources: Recognizing the sovereign rights of States over their natural resources, the authority to determine access to genetic resources rests with the national governments and is subject to national legislation.

Other key international mechanisms include CITES, the Ramsar Convention on Wetlands, the United Nations Convention on the Law of the Sea, UNESCO and its International Oceanic Convention. Recently, the World Summit on Sustainable Development,1 the Millennium Development Goals,2 and the Millennium Ecosystem

1 http://www.unep.fr/outreach/wssd/postjoburg/wssdoutcomes.htm
2 http://www.un.org/millenniumgoals/
Assessment\(^3\) have introduced broad goals into the international development arena. Specific goals have been identified in high priority areas such as Africa.\(^4\)

4. DRIVERS IN THE INTERNATIONAL POLICY SECTOR
The following trends may act as drivers of change in the use and value of aquatic genetic biodiversity and how international agencies deal with the changes.

**Trends in human, economic and biodiversity resources**
The production from capture fisheries has levelled and significant increases in production are expected to come primarily from aquaculture (Figure 1). Of the world’s major marine fisheries, the percentage of over-exploited, depleted or recovering stocks has increased from about 10% to 28% from 1974 to 2003. During this time under to moderately exploited stocks have decreased from about 40% to 24%, and fully exploited stocks have remained fairly constant at 50% (FAO, 2004).

Aquaculture is the fastest growing food producing sector with an average rate of increase of about 9% over the last two decades (FAO, 2004). Much of this growth has been in developing countries. Today, nearly one of every two fish consumed with be farm-raised (FAO, 2006a). It is further expected that per capita consumption of fish will increase to about 16kg/yr by 2015 (FAO, 2004). With an ever growing human population, fishery production will need to increase to meet these expectations. In response, intensification of farming systems, exploration of new areas and improvement of fishery management are being employed. Intensification has also involved genetic

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\(^3\) [http://www.eco-index.org/search/pdfs/millenium_ecosystem_assessment.pdf](http://www.eco-index.org/search/pdfs/millenium_ecosystem_assessment.pdf)

alteration of species to be consumed. This has led in some cases to a fear of products derived from modern biotechnology.

**Trends in technology**
At present there is substantial technical ability to identify, characterize and manipulate genetic biodiversity, although it is often not used or is expensive to use. Technology exists that allows genes to be transferred across taxonomic kingdoms, e.g. fish anti-freeze protein genes that have been inserted into strawberries; micro-satellite markers can trace family pedigrees or identify stock structure in wild populations (Liu, 2007), and the products of individual genes can be identified and crafted to meet certain needs. Improvements in technology have increased our ability to perform these manipulations and analyses and have lowered the cost of doing so. Although much of this technology is in developed countries, numerous developing countries have this capacity and many others are seeking it.

**Societal trends**
In response to the above increase capacity in technology, there has been increased attention given by consumers, non-governmental organizations, and other interests groups to human health safety, environmental safety and ethical concerns associated with genetic manipulation and consumption of fish and fish products. Many of the human health concerns stem from highly publicized aspects of plant genetic engineering where the products of the modification are toxins or resistance to toxins, e.g. herbicide resistant soybeans or Bt-cotton.

**From common heritage to sovereign rights**
With the signing of the CBD and the CCRF, the international community acknowledged the value of biological diversity and genetic resources in helping improve the human condition. Whereas previously genetic resources were considered to be the “heritage of mankind”, these new instruments now recognize the sovereign rights of States to manage their own resources and control access to them (CBD, 1994).

**Recognizing stocks and strains**
Responsible management of aquatic genetic resources will require information on stocks, strains, and important breeds of aquatic organisms (Grant, 2007; Pullin, 2007; Smith, 2007). Important stocks of marine species have been defined and assessed as to their status, i.e. depleted, recovering, sustainably harvested, over-fished. Some National Governments are granting species status to sub-species and stocks as in the United States of America where the government affords protection to endangered runs of Pacific salmon as species under the USA Endangered Species Act.5 Breeding centers in Eastern Europe maintain detailed information on strains of common carp (Bakos and Gorda, 2001), and registries of common aquaculture species exist in the United States of America;6 these are exceptions however, to the general lack of information below the species level.

**From simple to complex issues**
Although basic information on aquatic genetic resources and biodiversity is extremely important and much work still needs to be done to assess their status and trends,
numerous international and donor agencies and conventions are now stressing the important role that these resources play in poverty reduction, human health, and ecosystem functions (see for example Toledo and Burlingame (2006) and references therein). The complex issues of poverty and livelihoods are becoming superimposed on the technical issues of genetics and biodiversity. The CBD, CG Centers and FAO are working to document not only the aquatic animal diversity found in rice fields, but also the key nutrients such as fatty acids, minerals (Toledo and Burlingame, 2006). While this trend is expected and reasonable, it puts added importance to accurate assessment of aquatic resources for food and aquaculture.

5. REFERENCES


1. SUMMARY

Genetic diversity encompasses three hierarchical levels: differences between species, differences among conspecific populations and genetic differences among individuals in a population. While the protection of each of these levels of genetic diversity is essential for achieving sustainable harvests, overfishing, habitat degradation and climate change generally overshadow concerns for genetic integrity. Capture fisheries for freshwater and diadromous species are marginally increasing globally, but capture fisheries for marine species have leveled or are declining. The demand for fishery products remains unabated and will increase as the economies of developing countries improve.

The continuing development of new molecular genetic tools provides high-resolution markers for assessing genetic population structure, for estimating demographic parameters and for providing insights into breeding biology. A growing body of population and evolutionary theory, and new statistical and computer procedures greatly assist in the interpretation of genetic data. Presently, genetic variables are generally not incorporated into ecological or economic models. Future models incorporating genetic data will be tailored to particular situations.

Fisheries in rivers and lakes are largely focused on species with naturally fragmented populations. These species are prone to inbreeding depression in small populations and to hybridizations with introduced divergent strains. Hence, genetic concerns are usually addressed under the framework of conservation biology and theory relating to inbreeding and unintentional hybridization.

Diadromous species support large commercial fisheries in the North Pacific and North Atlantic. These species are especially vulnerable to ecological disturbances because of their complex life-history cycle, which spans freshwater and marine habitats. The loss of between-population genetic diversity through population extinctions in some species is especially acute in areas of human development. The failures of numerous transplanting programmes for many species indicate that local populations are adapted to particular habitats and seasonal events and cannot be easily moved to other habitats.

In the marine realm, the greatest genetic threats appear to be the extinction of genetically unique subpopulations and loss of genetic diversity through declines in abundance by overfishing and climate change. For species or stocks supplemented with cultured individuals, genetic swamping with artificially propagated individuals can reduce the fitness of wild populations.

Numerous international conventions and agreements recognize the importance of maintaining biological diversity, but generally treat genetic diversity indirectly as a component of biodiversity. Four steps provide a framework for conserving genetic
diversity: 1) identification of objectives, 2) assessment of genetic risk, 3) identification of reference points and 4) monitoring of progress toward objectives.

2. INTRODUCTION

Biological diversity encompasses three components: ‘the variety of living forms, ecological roles they perform and the genetic diversity they contain’ (Wilcox, 1984). Capture fisheries are faced with several problems that tend to erode these fundamental components of diversity. The most important problem in many environments is overfishing (Pauly et al., 1998, 2003; Allan et al., 2005), but habitat changes from human development, pollution and physical degradation from trawling are also substantial. The increasing demand for fish and weak enforcement of fishery regulations in many regions have led to serious depletions of once abundant stocks. These problems are especially acute in coastal and estuarine areas close to human development. In addition to these direct human impacts on wild populations, natural (North Atlantic Oscillation, Pacific Decadal Oscillation) and induced (climate warming from greenhouse gases) shifts in climate greatly influence the abundances of local populations (Attrill and Power, 2002; Benson and Trites, 2002).

The chief focus for achieving sustainable harvests of capture fisheries has been on the preservation of species abundances and ecosystems with little attention given to intraspecific diversity (Ryman et al., 1995). The reasons for this are twofold. First, management policies are heavily influenced by economic demand and the sustainable use of particular species. Second, the task of characterizing intraspecific diversity for each species is immense and often beyond the will or research capabilities of management agencies, especially those in developing countries. However, the maintenance of intraspecific genetic diversity may be key to preventing species extinctions (e.g., Ehrlich, 1988). The erosion of intraspecific diversity is not limited to small and geographically isolated populations but can also occur in seemingly abundant marine species.

Genetic resources can be viewed as genetic differences at three hierarchical levels of organization: 1) species, 2) populations and 3) individuals. At the highest level, species consist of populations that are reproductively isolated from populations of other species. Genetic isolation occurs because of geographic (allopatric) or behavioural isolation and, together with local adaptation, leads to the appearance of novel genetic traits (Otte and Endler, 1989). Hence, each species harbours a unique set of genetic material. Biologists agree that the process of speciation usually occurs on timescales of several hundreds of thousands of years. However, once species are lost, the fossil record indicates that several million years are required for species diversity to recover (Briggs, 1995).

At the population level of organization, the identification of discrete stocks has been a major theme in fisheries research. The definition of a stock can vary, as the motivations of fishery managers may be influenced by political, economical or biological mandates (Carvalho and Hauser, 1994). As a result, management boundaries are sometimes set at national borders because of issues of jurisdiction, even though a biological perspective may be of far greater importance in promoting the viability of a stock. The problem of managing “straddling stocks” is of particular importance for many highly mobile marine species (Meltzer, 1994).

Finally, the largest store of genetic variability in most species exists as genetic differences among individuals within a population. This variability arises from the physical assortment of genes among offspring during reproduction. Of great importance for the conservation of this genetic variability is the theoretical concept of effective population size, which is usually much smaller than census size. Both theory and empirical results show that the loss of genetic variability is greater in small populations than in large populations. Hence, the goal of preserving genetic variability
in a population coincides with the goal of maintaining large ecologically sound natural populations.

In agriculture, the problem of conserving genetic diversity has been largely framed as the preservation of domesticated plant cultivars and animal breeds, which have adapted to local environments over thousands of years of selective breeding. Technical advances have led to a greater availability of cheaper grains, and this has produced a shift from pastoral grazing to more capital-intensive methods of farming. Intensive farming methods are more productive and more predictable than traditional methods of farming. Consequently, farmers have abandoned many indigenous breeds, and this shift has led to the loss of genetic diversity. Much less attention has been directed toward the conservation of genetic resources in natural, free ranging capture species. The development of domesticated breeding lines for aquatic organisms is still in its infancy and depends on the availability of wild strains to a much greater degree than does the present-day development of breeds of plants and animals for agriculture.

The chief goal of this study paper is to survey the status of genetic resources in freshwater and marine capture fisheries and to develop an argument for conserving genetic resources in these species. These arguments parallel those developed for the conservation of plant and animal genetic resources. A second goal is to outline trends in the development of these methodologies and the concepts used to manage genetic variability in capture fisheries. The methodologies used to describe genetic variability and to assess its value in inland and marine capture species differ somewhat from those used to assess genetic resources in domesticated plants and animals. A third goal is to summarize institutional mandates focused on preserving genetic diversity and to present a framework of action for conserving genetic diversity.

2.1 Why conserve genetic diversity?
Several arguments have been developed to support the notion that the conservation of genetic resources is important in various settings. Biological and normative justifications for conserving genetic diversity are:

1. to ensure the future adaptability of natural populations;
2. to preserve life-history, behavioural and morphological traits that ensure sustainable fisheries;
3. to promote the use of genetic resources in commerce and medicine; and
4. to conserve genetic diversity for cultural reasons.

Although these arguments have been developed for agricultural resources, they are a starting point for developing analogous arguments for the conservation of diversity among and within species supporting capture fisheries.

In agriculture, indigenous breeds have value for the creation of new breeds, even though individually they may not be of high economic value (Mendelsohn, 2003). Locally adapted breeds, for example, may harbour genes that promote disease resistance, which may have been lost in highly selected production strains. Other arguments focus on societal choices. A society may be willing to maintain economically inferior breeds, because these breeds may be part of a local landscape that is valued by society, or because society finds value in maintaining historical activities and traditional livelihoods. The decline of indigenous breeds is often tied to biological and environmental conservation issues in developing countries.

Much less attention has been given to evaluating the importance of genetic resources in species supporting capture fisheries. The chief reason is that little is known about the genetic components of production in wild populations. These populations lack the recorded breeding histories that are maintained for plant cultivars and domesticated livestock. Although many inland and some diadromous species can be bred in captivity, only a few marine species have been bred in captivity. A compelling reason for conserving genetic diversity in wild populations is to provide a large base for
developing strains for aquaculture. Wild populations of plants and animals are now no longer used to a large extent to develop new agricultural strains. However, the development of strains for aquaculture is ongoing and depends on the availability of genetically diverse wild populations.

2.2 Trends in capture fisheries production

Inland and marine capture species together make up the bulk of fishery products, although production from aquaculture is increasing rapidly. Fisheries provided about 140 million tonnes of food and fish products in 2001. Most of this production comes from marine waters (about 85 million tonnes; 59.8%) and almost half consists of small pelagic fishes. The remaining capture production comes from inland waters (8.7 million tonnes; 6.1%) (FAO, 2003). A growing amount of production comes from marine and freshwater aquaculture (48.4 million tonnes; 34.1%). The size of the marine capture fishery has leveled in the last few years and may be declining (Pauly et al., 2003), while inland fisheries have been relatively stable, or marginally increasing (Figure 1). Inland capture fisheries are largest in Asia (5.8 million tonnes) and Africa (2.1 million tonnes), with important fisheries also in Europe (0.3 million tonnes), South America (0.3 million tonnes), North America (0.2 million tonnes) and Oceania (0.02 million tonnes). About 7 million tonnes (80% of inland fisheries) are produced in countries with low average incomes and food deficits. Inland capture fishery production is the sole source of fish in many of these countries.

2.3 Trends in demand for fishery products

Trends in the consumption of fish suggest continued increases in the demand for fish. A sample of 132 nations indicates that the consumption of fish is greatest in countries with high standards of living, as measured by per capita gross domestic product (York and Gossard, 2004). However, demand differs among regions and among nations. Several developing countries have high fish consumption, including Bangladesh, Cambodia and China (FAO, 2003). The demands for fish products in the nations of Africa, the Middle East and western countries of North and South America and Europe are similar (Figure 2). The largest rate of increase occurs in Asian countries, because of the traditional emphasis on fish consumption, population increases and economic

![FIGURE 1](image-url)

**FIGURE 1**
Marine, inland capture fishery production, and aquaculture production (FAO)
improvement. The shift of rural populations into cities, which often accompanies economic development, also leads to the increased consumption of fish. These trends indicate that the demand for fish will increase globally, but will increase most in Asian countries as they develop economically.

3. USE OF MOLECULAR MARKETS TO SURVEY GENETIC RESOURCES

The use of molecular genetic markers to survey genetic variability and to infer population processes has advanced on two fronts in recent years. New technologies have been developed to assay DNA polymorphisms directly, and these methods have produced a range of DNA markers with complementary characteristics to address various questions (Annex 1). These advances provide a means of generating large amounts of data as a basis for statistically testing research and management hypotheses.

3.1 Applications

Assessments of genetic variability in species in capture fisheries are important for several reasons. Molecular genetic markers occur naturally, are inherited in a predictable way, provide a basis for rigorous statistical analysis, and thus are ideally suited to assessing genetic variability in wild populations. Molecular markers have been used to discover morphologically cryptic species (Shaklee and Tamaru, 1981; Knowlton, 1993; Bernardi and Goswami, 1997; Knowlton et al., 1997), define population boundaries (Ruzzante et al., 1998; Waples, 1995), estimate population components in areas of population mixing (Hansen et al., 2001; Nielsen et al., 2001; Hauser et al., 2006) or origins of juveniles during life-history migrations (Teel et al., 2003; Bowen et al., 2006). Box 1 illustrates the use of mitochondrial DNA markers to infer migration patterns of juveniles of endangered hawksbill sea turtles (Eretmochelys imbricata).

One promising use of molecular data is to estimate population parameters such as population size (Nunnery and Elam, 1994; Bagley et al., 1999; Turner et al., 2002) (Annex 1) or to reconstruct demographic histories of population growth or
Molecular genetic data have been especially useful for estimating kinship among individuals in natural populations (Bernatchez and Duchesne, 2000; Bentzen et al., 2001; Garant et al., 2001; Banks et al., 2003), for measuring reproductive success (Fiumera et al., 2002) or for forensic identifications (Birstein et al., 2000).

The development of high-resolution population markers provides a means of testing models of population structure. Populations of inland species are largely isolated from one another by terrestrial barriers to movement and the construction of bottlenecks in population size (e.g. Luikhart et al., 1998a). An understanding of responses to past environmental or climate disturbance can give clues to how populations might respond to future challenges. Molecular genetic data have been especially useful for estimating kinship among individuals in natural populations (Bernatchez and Duchesne, 2000; Bentzen et al., 2001; Garant et al., 2001; Banks et al., 2003), for measuring reproductive success (Fiumera et al., 2002) or for forensic identifications (Birstein et al., 2000).

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The development of high-resolution population markers provides a means of testing models of population structure. Populations of inland species are largely isolated from one another by terrestrial barriers to movement and the construction of

**BOX 1**

**Genetic mixed-stock analysis of Caribbean juvenile hawksbill sea turtles, a CITES listed species. (Bowen et al., 2006)**

Genetic stock identification has been especially useful in the management of species that are harvested in areas of stock mixing. In these areas, less abundant stocks may be threatened with overfishing. The method was developed to estimate the proportions of component stocks in harvests of Pacific salmon as they returned to rivers to spawn (Grant et al., 1980), but has proved useful for other species.

Hawksbill turtles (*Eretmochelys imbricata*) are specialized sponge feeders, which migrate between nesting beaches and feeding habitats on tropical reefs. The colourful “tortoiseshell” scutes of this species are especially valued in the production of artisan products. Harvests of this species have brought it close to extinction. A pressing question has been the extent that harvests of juveniles on feeding grounds influence spawning site abundances in other areas, a perennial problem in ‘straddling stock’ species. The migration biology of juveniles is largely unknown because the physical tagging of nestlings is impossible. A survey of mtDNA variability showed strong haplotype frequency differences among female nesting sites that could be used to identify the origins of juveniles on shallow reefs (Bowen et al., 2006). Bayesian estimates of the origins of 629 juveniles from seven feeding congregations demonstrated that juveniles tend to return to feeding areas close to their birth sites. A significant correlation appeared between the percentage contribution to a feeding area and the distance from the contributing population (Figure). The magnitude of these distances indicates that harvests in one part of the Caribbean will impact nesting sites throughout the region.
population models is straightforward. These populations generally act as collections of subpopulations (a metapopulation), in which subpopulations are tied to each other by various levels of gene flow. Local extinctions and colonizations also appear to be a general feature of inland (e.g., Bernatchez and Wilson, 1998; Lafferty et al., 1999).

The structures of marine populations, on the other hand, are less well known. As expected, near shore species with both limited larval and adult dispersal (or homing behaviour to spawning areas) tend to have subdivided population structures (McQuinn, 1997; Robichaud and Rose, 2001). However, many species in capture fisheries have high dispersal abilities and occur in oceanic areas without firm barriers to movement. These species tend to show much less genetic population structure with populations occupying much larger areas than do populations of inland species (see below). How these populations are structured is of considerable importance to their management in capture fisheries. Surveys of molecular population markers continue to be important for testing the various models of population interconnection and structure.

These applications generally assume that molecular markers are not directly influenced natural selection and that the distributions of the markers reflect such parameters as effective population size and gene flow. However, genetic diversity itself is also an important component of ecological and evolutionary health of a species. A rapidly growing field of research focuses on the development of molecular markers linked to quantitative trait loci (genes that affect the ecological fitness of individuals). "Genomic" methods are used to survey portions of the genome directly influenced by selection (Reid et al., 2005; Slate, 2005), to monitor genotoxic pollutants (Newton et al., 2004, Rockett and Dix, 1999) and to study the effects of hybridization (Dowling and Childs, 1992; Rhymer and Simberloff, 1996) and population crashes on gene organization (Luikart et al., 1998a, b; Garza and Williamson, 2001).

Market or production traits, including growth rate, flesh characteristics and disease resistance, are generally influenced by the actions of several genes, which can be evaluated only by breeding experiments (e.g., Law, 2000). In addition to experimental breeding manipulations, information on breeding lines and pedigree analysis form the basis for evaluating genetic resources in domestic livestock and agricultural plants. However, laboratory experiments on most populations targeted in capture fisheries are not possible, so alternative methods are required to assess the genetic status of natural populations.

### 3.2 Statistical analysis

The development of statistical methodologies and computer programmes has kept pace with laboratory progress in providing a means of analyzing genetic data (Annex 2). A variety of statistics can be used to assess genetic diversity within and among populations. Genetic data for a sample of individuals can provide information about genetic diversity within and among populations (Nei, 1987; Hedrick, 2005) and can be used to infer phylogenetic relationships among species (Felsenstein, 2004). Three measures of diversity are widely used in conservation and population studies (Box 2). The first, average heterozygosity, $H$, (also called gene diversity) measures the level of genetic variability in a population and is routinely estimated with allozyme and microsatellite DNA data (Nei, 1987). Nucleotide diversity, $\Theta_\pi$, extends the concept of gene diversity by adding a measure of sequence divergence between haplotypes. These statistics can be used to detect the erosion of genetic diversity from historical reductions in population size. A third statistic, $F_{ST}$, measures diversity among subpopulations in a species.
BOX 2
Statistics used to measure genetic diversity within and among populations

Average heterozygosity or gene diversity: Average heterozygosity can be estimated in two ways. The first way is to count the number of heterozygous individuals in a sample of diploid genes. This is known as observed heterozygosity, $H_e$. These counts are usually presented as a proportion of all genotypes. A second way of estimating heterozygosity is to assume that the sample of genes from a population does not deviate significantly from Hardy-Weinberg proportions and calculate the proportion of expected heterozygotes from gene frequencies. For a single locus expected heterozygosity, $h$, can be calculated as:

$$h = 1 - \sum p_i^2$$

where $p_i$ is the frequency of the $i$th allele in a sample. This formula is used to also estimate gene diversities for haplotypic loci such as mitochondrial DNA in animals or plastid DNA in plants. When data for a sample of several loci are available, such as for allozyme and microsatellite DNA data, heterozygosities are averaged,

$$H = \frac{\sum h}{R},$$

Where $R$ is the number of loci sampled. Average heterozygosities estimated from allozyme data usually also include monomorphic loci and are taken as an estimate of genome wide variation when samples sizes of loci are large ($R > 20$). Average heterozygosities based on microsatellite DNA usually only include polymorphic loci and are, therefore, not comparable with allozyme heterozygosities.

Nucleotide diversity: Sequences of DNA provide a basis for estimating divergences between alleles, which is not possible for allozyme or microsatellite data. The amount of sequence divergence between haplotypes in a sample provide information about the age and historical size of a population. Other variable equal, older populations are expected to accumulate more mutations and show larger divergences between haplotypes. One the other hand, larger populations of the same age are also expected to accumulate a greater number of mutations. The loss of low frequency haplotypes in a large population is less than in small populations because the loss of haplotypes through genetic drift is less. These haplotypes, however, are expected to be closely related to each other. These characteristics form the basis for estimating several demographic parameters of populations (Rogers and Harpending, 1992).

Nucleotide diversity can be estimated from the average number of nucleotide differences between haplotypic sequences, $\pi$. The number of nucleotide differences per nucleotide site, $d_{xy}$, is used to account for differences in the lengths of sequences in different studies. Nucleotide diversity, $\Theta_\pi$, is the sum of the product of divergences between haplotypes and the frequencies of haplotypes

$$\Theta_\pi = \sum \sum d_{xy} p_x p_y,$$

Where $p_x$ and $p_y$ are frequencies of haplotypes in a sample.
**Box 2 (cont.)**

$F_{ST}$ This statistic is the standardized variance of gene frequencies among populations and is estimated by

$$F_{ST} = \frac{\text{var}(p)}{p(1 - p)}$$

where $p(1 - p)$ is the binomial variance. This statistic ranges from 0.0, indicating identical gene frequencies between populations, to 1.0, indicating fixed gene frequency differences between populations. $F_{ST}$ values are usually averaged over loci when data for several loci (allozymes and microsatellites) are available. The co-distribution of $H$ and $F_{ST}$ can be used to test for the effects of natural selection on gene frequency divergence between populations (Beaumont and Nichols, 1996).

The maximum value of $F_{ST}$ is limited by high heterozygosities, as are commonly found for microsatellite loci. The upper limit of $F_{ST}$ is $(1 - H_s)$, where $H_s$ is the average within subpopulation heterozygosity (Hedrick, 1999). Hedrick (2005), therefore, suggests that $F_{ST}$ be modified

$$F'_{ST} = \frac{F_{ST} (1 + H_s)}{1 - H_s}$$

These adjusted values provide a better estimate of differentiation among populations when estimate with microsatellite DNA.

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**4. Patterns of Genetic Viability in Natural Populations**

Species generally are divided into a few to numerous populations depending on the dispersal ability of individuals and on the availability of dispersal corridors. At one extreme, some marine fishes with highly mobile adults and with unimpeded movements of planktonic eggs and larvae, consist of only a single global population (Figure 3). At the other extreme, some species are highly subdivided into genetically unique subpopulations because of physical barriers to movement, or because of strong natal and site spawning fidelity. As a consequence, many marine species, such as tunas, tend to be "population poor", while inland and anadromous species tend to be "population rich" with numerous small populations. How populations react to physical and biological variables is a subject of ongoing debate (Sinclair, 1988; MacCall, 1990; Sherman et al., 1993).

Various isolating mechanisms produce different levels of population subdivision among inland, anadromous and marine populations (Table 1). Although the relative amounts of allozyme and microsatellite DNA gene diversity are similar among these groups, how this diversity is partitioned among populations differs among groups. The largest amount of genetic subdivision appears among conspecific populations of freshwater species (mean $F_{ST} = 0.222$, median $F_{ST} = 0.144$), because of the physical isolation of lake and riverine habitats. Anadromous species (salmonids) show large amounts of population subdivision (mean $F_{ST} = 0.108$, median $F_{ST} = 0.081$). In this group, subdivisions reflect not only geographic isolation between freshwater spawning sites, but also homing to natal spawning sites. Populations of marine fishes show the least amount of genetic subdivision (mean $F_{ST} = 0.062$, median $F_{ST} = 0.020$), because of fewer restrictions to the movement of eggs, larvae and adults in marine waters. These statistics have been used to infer the number of migrants between populations each generation. However, the models used to make these estimates are over-simplifications...
A. Independently, self-sustaining population that do not exchange individuals. These populations are expected to show substantial life history, demographic, morphological or genetic differences.

B. Partially isolated populations with some geographic overlap or exchange of individuals, or both. These populations may show small life history, demographic, morphological or genetic differences. Genetic differences are often used to define genetic stocks. Theoretically genetic differences appear between populations over long periods only when migration is limited to 1-5 individuals each generation.

C. Substantial geographical overlap or mixing of individuals. Although life history or demographic differences may still appear between populations, genetic differences are not expected to appear. Genetic methods are incapable of detecting these populations, however, from a fishery management perspective each population may still merit recognition.

D. Panmixia. Only a single population exists with individuals (or gametes) freely moving between areas.

TABLE 1
Components of gene diversity in freshwater, anadromous and marine fishes (summarized from Ward et al., 1994 and Waples, 1998). \( H_T \) is the total amount of genetic diversity in a species, and \( F_{ST} \) is the variance of allozyme frequencies among subpopulations. \( H \) for microsatellite DNA is based on a single or only a few populations (DeWoody and Avise, 2000)

<table>
<thead>
<tr>
<th>Group</th>
<th>Number of Species</th>
<th>Mean number of populations in sample</th>
<th>( H_T )</th>
<th>( F_{ST} )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Average</td>
</tr>
<tr>
<td>Allozymes</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Freshwater</td>
<td>49</td>
<td>5.9</td>
<td>0.062</td>
<td>0.222</td>
</tr>
<tr>
<td>Anadromous</td>
<td>7</td>
<td>13.1</td>
<td>0.057</td>
<td>0.108</td>
</tr>
<tr>
<td>Marine</td>
<td>57</td>
<td>6.4</td>
<td>0.064</td>
<td>0.062</td>
</tr>
<tr>
<td>Microsatellite DNA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Freshwater</td>
<td>13</td>
<td></td>
<td>0.54</td>
<td></td>
</tr>
<tr>
<td>Anadromous</td>
<td>7</td>
<td></td>
<td>0.68</td>
<td></td>
</tr>
<tr>
<td>Marine</td>
<td>12</td>
<td></td>
<td>0.66</td>
<td></td>
</tr>
</tbody>
</table>

of real populations. Hence, these estimates should be used cautiously (Bossart and Prowell, 1998; Whitlock and McCauley, 1999).

4.1 Inland species
Even though inland capture fisheries are much smaller than marine capture fisheries, freshwater fisheries are an important source of protein in many countries. About 7 million tonnes (80 % of inland fisheries) are produced in countries with low average incomes and food deficits (FAO, 2003). Fisheries are the sole source of animal protein in many of these countries. Freshwater habitats are much more fragmented than marine habitats and experience greater seasonal extremes in temperature and desiccation. Hence, freshwater populations of fishes are expected to be smaller in general, show greater fluctuations in abundance and be genetically more subdivided
than marine fishes. The problems facing populations of freshwater species tend to be addressed largely by principles in conservation biology, rather than in fishery management. Even though the problems facing inland aquatic species are most severe in developing countries, aquatic species in developed countries have best been studied with genetic methods.

4.1.1 Habitat degradation and genetic population structure
The greatest problem facing inland aquatic species is habitat degradation from human activities. Aquatic habitats are often modified by deforestation and watershed erosion, pesticides and agricultural and industrial run-off. Canalization of streams for agriculture, or direct use by humans, destroys riparian zones and impairs natural ecosystem processes that maintain water quality and produce food for aquatic species. Habitat degradation can have important genetic consequences for aquatic populations.

1. Habitat-related reductions in population size inevitably lead to the loss of genetic diversity and often to inbreeding depression. Genetic diversity can decline rapidly in small populations through random genetic drift (Crow and Kimura, 1970). The detrimental effects of inbreeding are well known from agriculture and aquaculture and inbreeding itself can lead to the demise of a population (e.g., Gall, 1987; Leberg and Vrijenhoek, 1994).

2. Habitat degradation often leads to habitat fragmentation and the loss of important connections between populations. The loss of habitats increases genetic isolation and reduces the possibility of genetic rescue of endangered populations and the probability of colonization of empty habitats in a metapopulation (Hanski and Gilpin, 1997).

4.1.2 Genetic risks from introduced species and non-native stocks
Another important threat to inland species is the introduction of non-native species, or of stocks of the same species adapted to different environments. Introductions can produce a variety of effects.

First, introductions of non-native species can lead to ecological imbalances. For example, the introduction of the predatory Nile perch (Lates niloticus) into the Great Lakes of Africa led to a population explosion that caused the extirpation of about 65% of Lake Victoria’s endemic cichlid fish species diversity (Witte et al., 1992; Goldschmidt et al., 1993). The large populations of Nile perch then supported a large fishery in Lake Victoria and the production of choice filets for European and Asian markets (Kitchell et al., 1997). In recent years, the burgeoning fishery has in fact reduced predation pressure on native species to about 10% of its 1970 levels and has allowed the rebound of some of the remaining cichlid species (Kitchell et al., 1997).

Second, fish are sometimes transferred to other areas out of their native range. If the transferred fish are genetically different from local fish, hybridization between the two may lead to outbreeding depression and the loss of fitness (e.g. Morizot et al., 1991; Carmichael et al., 1993). Box 3 gives a case history of stock supplementation and mixing of two subspecies of largemouth bass across North America. Alternatively, if the introduced individuals are competitively superior to local individuals, local native populations may be hybridized to extinction.

Third, introduced individuals of the same species may be genetically compatible with local populations, a condition assumed in most stock supplementation programs and hatchery operations. Great caution, however, is needed to ensure that artificially propagated individuals have not also been genetically modified by adaptation to hatchery conditions. Selection in hatcheries can potentially occur with some feeding methods, the selection of broodstock, or other procedures that modify behaviour. Supplementation of wild populations by hatchery reared individuals can potentially lead to genetic ‘swamping’ and the loss of genetic diversity in wild populations, even
4.2 Diadromous species

Diadromous fishes include species that spawn in either fresh or marine waters, but spend part of their life cycle in the other habitat. Diadromous species exhibit several life history traits that make them vulnerable to extinction (Jonsson et al., 1999) (Box 4). About 18% of diadromous fishes are considered to be endangered, threatened, rare or vulnerable, whereas only about 5% of fish species in general are considered to be of conservation concern (Barbault and Sastrapradja, 1995; McDowall, 1999).

4.2.1 Catadromous species

These fishes spawn in marine waters, but migrate into fresh or brackish water. The best studied of these species are North American (Anguilla rostrata), European (Anguilla
anguilla), Asian (Anuillla japonica) and shortfinned (Anguilla australis) eels, which spawn in the marine waters, but mature in rivers and lakes. In North America, allozyme markers showed significant differences among river populations (e.g. Williams et al., 1973), whereas mtDNA markers indicated a lack of geographic differentiation (Avise et al., 1986; Lintas et al., 1998). Recent studies with high-resolution molecular markers have detected weak, but significant, differences among freshwater populations of European eels (Daemen et al., 2001; Maes and Volckaert, 2002; Wirth and Bernatchez, 2003), but not among North American populations (Wirth and Bernatchez, 2001). Species consisting of a single large breeding population are particularly vulnerable to environmental changes and require international cooperation in their conservation and management.

4.2.2 Anadromous species

Anadromous fishes, on the other hand, spawn in freshwater, but mature in marine waters before returning to freshwater to spawn. The problems facing these species are particularly severe because, in addition to harvest mortality, individuals are tied to aquatic habitats that are often heavily impacted by human activities. One group of special concern includes anadromous and freshwater resident species of sturgeons, which exhibit many of the life history traits predisposing species to extinction (Williot et al., 2002) (Box 4). They occupy different habitats during their life cycle, are large, long-lived, slow growing and late maturing. Habitat degradation, dams and exploitation for caviar have produced alarming population declines (Birstein, 1993). Genetic studies of sturgeons indicate complex population structures (Doukakis et al., 1999; Campton et al., 2000; Wirgin et al., 2000) and confused taxonomies (Phelps and Allendorf, 1983; Birstein et al., 2000; Birstein et al., 2002; Krieger et al., 2000).

Anadromous salmonids, which are distributed across temperate regions of the Northern Hemisphere, are also vulnerable to many of the same threats facing sturgeon (Nehlsen et al., 1991). Considerable effort has been spent on the life history and genetic analyses of these species, because these fishes support substantial commercial, traditional or recreational fisheries. A variety of genetic population structures appear in salmonids that reflect ecological and life history differences, not only among species, but also among some stocks of the same species. For example, Waples et al. (2001) found that among seven species of Pacific salmon in the genus Oncorhynchus a strong correlation appeared between ecological and life history diversity. Species inhabiting a greater number of ecological provinces tended to exhibit a greater number of life history types as evidenced by degree of anadromy, spawning run timing, time to adult maturity (marine phase duration) and juvenile freshwater residence time (Figure 4). This correlation reflects the direct influence of environmental factors such as temperature and food availability on the expression of life history traits.

An understanding of the nature of adaptive traits is of fundamental importance in the conservation and management of fishery resources. Both rate and mechanism determine the extent to which life history diversity and diversity generating eco-processes should be conserved. Adaptations in many salmonids occur rapidly (Hendry, 2001; Koskinen et al., 2002) and over short distances (Taylor, 1991). Life history characters shifted in only a few decades after introductions of Chinook salmon (Oncorhynchus tshawytscha) to New Zealand (Quinn et al., 2000) and into the North American Great Lakes (Kwain and Thomas, 1984). Rapid rates of life history diversification on contemporary time scales have also been documented in sockeye salmon (Oncorhynchus nerka) (Hendry 2001) and grayling (Thymallus thymallus) (Koskinen et al., 2002). The failures of many stock transfers of salmon between rivers and streams along the west coast of North America indicate
BOX 4

Population or biological traits that predispose stocks or species to depletion and extinction. Declines in effective population sizes can lead to the loss of genetic diversity

1. **Slow growing and long lived:** Species with these traits are vulnerable to the effects of overfishing because standing biomass after harvest is replaced very slowly. These traits are often associated with large body size, late maturity and small numbers of offspring.

2. **Several years to reproductive maturity:** Many species of fish have market value before they reach reproductive maturity. Unless a significant number of individuals are allowed to reproduce the viability of a stock is greatly reduced. Another component of this problem is that older, larger individuals often have reproductive potentials far larger than younger, mature individuals.

3. **Few offspring per year:** Producing only a few offspring per year is part of a continuum of reproductive strategies. Many fish and invertebrates produce millions of eggs with little or no parental care of larvae. Even though the probability of survival to maturity for individual eggs is very small, at least some of larvae are expected to survive. Other species invest more parental care by producing larger, but energetically more costly eggs, or by guarding offspring. These latter species are most at risk from the effects of overfishing, because fewer offspring are produced.

4. **Large body size:** Species with large body sizes are in jeopardy for two reasons. 
   1) Large, conspicuous animals may be easier to find and harvest than small animals. Visibility is especially detrimental when these species inhabit confined embayments and estuaries. 
   2) Large animals are inevitably at the top of the food chain and are particularly sensitive to shifts in abundance of species in the food web. Species with large body size are often slow growing, produce few offspring annually and consist of few individuals.

5. **Small natural population numbers:** These species are at particular risk when a large part of their habitat is degraded or destroyed. When core populations become depleted, recovery is hindered by reductions in the number of reproductively active individuals. These species may also be vulnerable to the loss of genetic variability and to such genetic effects as inbreeding depression.

6. **Live in confined habitats:** Species inhabiting confined spaces, such as lakes, estuaries or coastal embayments are much easier to capture than similar species inhabiting the open ocean. Many confined habitats are also associated with human activities, thus increasing the exposure to fishing and habitat changes.

7. **Specialized habitat or life history requirements:** Species with special requirements are at particular risk when only a few suitable habitats are available, or when populations of suitable prey species have been reduced. Species using rivers as migratory pathways may be at risk from the construction of dams and shoreline development. Other species with specialized diets may be at risk when particular items of food are no longer available.
a general lack of ecological inter-changeability between most subpopulations (Utter, 2004).

4.3 Coastal marine species
Most of the World’s capture fisheries focus on marine species, some of which support annual harvests of several million tonnes. Species supporting the largest harvests generally occur over the continental shelf in areas with high levels of productivity driven by upwelling. Nutrient rich areas in the eastern boundary currents of North and southern Africa, and North and South America, for example, support large fisheries of pelagic fishes, including hakes, mackerel, anchovies and sardines. Even though many species of marine fishes occur in very large populations, the combination of overfishing and climate change make them susceptible to extinction (Musick et al., 2000; Myers and Ottensmeyer, 2005).

4.3.1 Genetic population structures of marine species
Most marine fishes and invertebrates are broadcast spawners and hence have large potentials for movement between areas by larval drift in currents. Additionally, adults of many species are capable of making long distance migrations. In contrast, adult homing to spawning areas, larval behaviour and hydrographic barriers to movement tend to isolate populations from one another, but not to the same degree as with freshwater fishes. The problem of unraveling demographic and genetic components of stock structure from gene frequency data is especially acute because of ill-defined geographical boundaries and decadal shifts in distributions. For example, early genetic studies of marine fishes indicated that they generally had moderate levels of gene diversity and little population subdivision, often over several hundred kilometers (e.g. Grant 1985; Mork et al., 1985). However, recent studies with high-resolution markers, such as microsatellite DNA and mtDNA, have revealed fine-scale spatial differences (e.g. Ruzzante et al., 1998) and unsuspected deep genetic lineages (e.g. Magoulas et al., 1996).
How populations of marine fishes are structure is a subject of some debate. A persistent problem has been the lack of models that satisfactorily incorporate both ecological and genetic concepts of populations (Annex 3). Ecological models generally assume that populations are highly adapted to local environmental conditions (Sinclair, 1988). The genetic prediction of this model is that species should consist of genetically differentiated, locally adapted populations. This prediction is borne out by genetic data for freshwater and riverine species, but not for many marine species. Other models postulate that contemporary levels of gene flow or historical range expansions and contractions (MacCall, 1990) imprint genetic gradients on populations (e.g. Lecomte et al., 2004). The different implications of these two models are important to formulating management policies and planning locations of marine protected areas.

Genetic estimates of gene flow are high in most marine species (Table 1), implying the movements of tens and hundreds of individuals between subpopulations. Mitochondrial DNA data appear to support the basin model for California anchovy (Lecomte et al., 2004), but support a mosaic model for European anchovy (Grant, 2005; Magoulas et al., 2006). However, finer-scale differences have been detected among populations that are not isolated by obvious physical or hydrographic barriers (Hedgecock et al., 1994; Ruzzante et al., 1999). This chaotic variability is likely due to large reproductive variances among families (Hedgecock, 1994), rather than to isolation or to adaptations to particular open-water habitats. The instability of marine waters on annual, decadal and millennial time scales likely prevents adaptations to specific areas. On a decadal scale, anchovy populations, for example, respond rapidly to small climate changes with range contractions and expansions (e.g. Cushing, 1982; Beare et al., 2004).

4.3.2 Effects of fishing on genetic variability

Populations of marine fishes, especially species supporting harvests of millions of tonnes, are generally thought to consist of large effective population sizes, and hence to be immune to the same genetic problems facing small populations of inland and anadromous species. Effective sizes of marine populations, however, may be much smaller than previously thought, because large fecundities can lead to large variances in family success. Only offspring spawned during a narrow window of oceanic conditions conducive to larval survival eventually recruit into the adult population (Hedgecock, 1994). As a result, the genetic effective size of a population may be orders of magnitude smaller than its census size (Nunnery and Elam, 1994; Bagley et al., 1999; Turner et al., 2002). Empirical evidence for this hypothesis, however, is mixed (Ruzzante et al., 1996; Herbinger et al., 1997; Li and Hedgecock, 1998). Nevertheless, available evidence indicates that fishing pressures can alter the genetic and demographic structures of seemingly very large marine populations.

The use of molecular genetic markers to estimate contemporary gene diversities is problematic since diversity is influenced by long-term rather than short-term effective population sizes. Long-term effective population size is the harmonic mean of populations each generation, and this mean is most influenced by small populations sizes. Abundances of most populations of marine species fluctuate on decadal and millennial time scales. For example, the analysis of fish scales in anaerobic sediments in the Santa Barbara Basin indicates large shifts in the abundances of anchovies and sardines over the last 2000 years before the onset of fisheries (Baumgartner et al., 1992). Spencer and Collier (1997) classified population fluctuation patterns of several marine fishes based on historical catch statistics. Three variables, coefficient of variation (CV) in abundance, variable around the long-term mean and temporal autocorrelation in abundance revealed five categories of population behaviour: 1) spasmodic, 2) high variability, 3) cyclic, 4) irregular and 5) steady state (Figure 5a). Allozyme data for many of the species in the Spencer and Collier (1997) study show a negative
relationship between heterozygosity and the CV of historical abundance (Figure 5b; Grant and Waples, 2000). These results indicate that even temporary reductions in population size can have a strong influence on long-term population size and, hence, on genetic diversity.

Genetic variability in a stock can also be lost through selection by capture methods, in addition to genetic drift and metapopulation dynamics. Effects of fishing on species are evidenced by shifts in the average sizes of individuals (e.g. Ricker, 1969, 1981; Bigler et al., 1996), changes in inherited life history parameters (e.g. Beacham, 1983a, b), reductions in average heterozygosity (e.g. Smith, 1994; Hauser et al., 2002) and by temporal shifts in gene frequencies (e.g. Lacson and Morizot, 1991). Directional selection can occur more rapidly in large populations than in small populations, because random drift, which tends to counter selection, is much less in large populations (Ryman et al., 1994).

5. GENETIC THREATS TO CAPTURE FISHERIES

Population size is a key variable for maintaining the genetic integrity of species in capture fisheries. Several ecological and genetic factors can converge to reduce population abundances. Most important for the marine environment has been overfishing by large industrial fleets. While external factors may play a role in some stock extinctions, intrinsic genetic factors can also be operating. One risk is the loss of genetic diversity, which declines at a rate that is inversely proportional to effective population size because of random genetic drift. The loss of genetic diversity can limit the ability of a population to adapt to changing environmental conditions and
detract from its economic value. Small populations face the risk of inbreeding (mating between close relatives), which increases homozygosity and, hence, the expression of deleterious, recessive genes.

5.1 Overfishing and habitat degradation

Overfishing has been implicated in the collapse of some fishery populations (Jackson et al., 2001; Allan et al., 2005). For example, Dulvy et al., 2003 documented the extinctions of 133 local, regional or global marine populations. Most of these extinctions could be attributed to overfishing (55%) or habitat loss (37%), while the remaining population declines appeared to be due to the effects of invasive species, climate change, pollution or disease. Global fisheries landings are continuing to decline at the rate of about 500 000 tonnes per year from a peak of 80-85 million tonnes in the late 1980s (Watson and Pauly, 2001). The effects of overfishing are not limited to large industrial fisheries. Small subsistence fisheries can also greatly influence species’ abundances (Jennings and Polunin, 1996; Friedlanner and De Martini, 2002).

These trends are likely to continue because of the increasing demand for fishery products and because of habitat degradation. As fisheries decline in productive waters over continental shelves, fishing is extended into deeper waters aided by the development of new technologies, such as satellite positioning and seafloor imaging. Marine species most vulnerable to stock depletions and extinction have large body sizes, long life spans, late maturities, low reproductive rates, limited geographical ranges, sporadic recruitment and adaptations to unique environments (islands and sea mounts) (Sadovy, 2001; Morato et al., 2006) (Box 4).

Once depressed, stocks may not recover for ecological and genetic reasons (Hutchings, 2005). For example, at very low abundances reproductive output falls off in some species (e.g. Shelton et al., 1999), and the removal of top predators may lead to dramatic shifts in ecosystem structure and food-web dynamics (Hansen et al., 1998; Scheffer et al., 2005). Ecosystem shifts resulting from ocean-climate changes may also retard the recovery of a depleted stock (e.g. Shelton et al., 2006). The reduction of population sizes by intense fishing also appears to have led to the loss of genetic diversity in some marine species (Smith, 1994; Hauser et al., 2002).

Declines in the abundances of natural stocks have stimulated aquaculture production. Although many countries have no alternatives, the aquaculture does not efficiently convert primary production into fishery products. Like the production of meat, which consumes about 40% of the world’s grain production to feed livestock (Harrison and Pearce, 2000), aquaculture requires large amounts of fish meal. Wastes from aquaculture often threaten nearby habitats and can severely degrade riverine and sensitive coastal ecosystems. Some aquacultural products are sometimes perceived to be inferior to products from capture fisheries (e.g. Hites et al., 2004; Senkowsky, 2004). Society, therefore, has a strong motivation to maintain wild stocks.

5.2 Genetic signatures of declining populations

The development of molecular methods provides an opportunity for identifying genetically distressed and declining populations. When populations become small, they are expected to lose genetic diversity. This loss is a central concern for declining populations threatened with extinction, as genetic factors can hasten extinction. For example, inbreeding depression is thought to be especially detrimental to the well being of small populations (Frankham, 1995; Hedrick and Kalinowski, 2000). Inbreeding between close relatives tends to increase the homozygosity of deleterious, recessive genes that are relatively harmless in the heterozygous condition. Low levels of gene diversity in themselves may not be universal distress signals, as many species with low gene diversities appear to be thriving after historical bottlenecks in
population size (Hoelzel, 1999). Conversely, moderate or high levels of gene diversity may mask genetic problems in a population. Hybridization, for example, may produce an artificially elevated level of gene diversity (Ferguson 1986; Leary et al., 1993).

Several biological and genetic indicators can be used to identify declining populations (Box 5). Molecular markers, for example, can be used to monitor levels of genetic

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**BOX 5**

**Genetic indicators of declining populations**

1. *Reduced gene diversity*. Genetic diversity can be lost in several ways. For a gene not under the influence of natural selection, the loss of gene diversity is inversely proportional to population size. Small populations lose diversity faster than large populations. Theory predicts that the loss of heterozygosity each generation from random genetic drift (reproductive sampling error) is 1/2N, where 2N is the number of gene copies in a population for a diploid gene. A recursion formula predicting the loss of gene diversity, \( b_t \), after \( t \) generations is

\[
b_t = b_0 (1 - 1/2N)^t
\]

where \( b_0 \) is the beginning level of gene diversity. Demonstrations of the loss of gene diversity in a population have to be made by comparison to un-fished populations of the same species and not to gene diversities in other species. Variability in gene diversity among species may be due to events on long evolutionary time scales and not to recent population events. Gene diversities can be measured with several molecular genetic markers, including allozymes, nuclear DNA (sequences or SNP polymorphisms) and mitochondrial DNA (RFLP or sequence polymorphisms).

2. *Changes in allelic or haplotype frequency distribution*. The Ewens’ (Ewens, 1972) sampling equation can be used to estimate an allele- or haplotype-frequency distribution from sample size and sample heterozygosity. This distribution is sensitive to the effects of population growth and decline and forms the basis of detecting recent bottlenecks in population size (Luikart et al., 1998a, b; Garza and Williamson, 2001).

3. *Genetic discontinuities among populations*. Geographical fragmentation resulting from population extinctions can lead to discontinuities in allelic or haplotypic frequencies. A demonstration of genetic population fragmentation, however, must be based on a comparison with populations inhabiting undisturbed environments.

4. *Altered phenotypic traits*. Low gene diversities, as measured by molecular markers, may not always detect populations in genetic distress. Selection on phenotypic traits, such as size at age, can be effective in large populations where random drift is unimportant. Shifts in average size for some capture species have been attributed to size-selectivity of fishing gear.

5. *Altered life-history traits*. The timing of fishing effort can alter the genetic profile of a population by eliminating some temporal components of diversity. For example, early spawning migrations in some species of Pacific salmon were eliminated because of fishing pressure on early returning adults.
diversity. In Atlantic salmon (*Salmo salar*), microsatellite DNA markers showed a decline in gene diversity in a contemporary population in Denmark, relative to gene diversity in archived scales from the same area (Nielsen *et al.*, 1997). Other genetic profiles can also be used to identify distressed populations, including the distributions of microsatellite DNA alleles (Garza and Williamson, 2001), the distributions of mtDNA frequency haplotypes (Tajima, 1989) and haplotype mismatch patterns (Rogers and Harpending, 1992). One problem in the application of some of these approaches is that the appearance of some genetic profiles often lags behind population declines, especially rapid declines. For example, Lavery *et al.*, 1996 found a mtDNA signature typical of an expanding population in a species that has declined in the past several decades.

### 5.3 Stock enhancement and supplementation

When capture populations decline, population enhancement and supplementation (the release of cultured individuals to boost wild population abundances) are sometimes used to attempt to rehabilitate wild stocks. Hatchery supplementations of salmonid populations have been practiced for several decades and provide lessons for other species (Utter, 2004). Although supplementation programmes for marine species have only recently been established, numerous projects are underway for fishes and invertebrates in several countries. Stock supplementations will likely increase as stocks continue to decline.

As aquaculture, mariculture and stock supplementation activities become more common, escapees and releases of cultured individuals will increase and potentially influence the genetic integrity of wild populations. The logic of supportive stock supplementation is to increase the survival of individuals in a hatchery without changing their genetic make up before release into the wild. However, the history of hatchery supplementation is filled with examples of genetic changes in cultured individuals, especially of salmonids (Reisenbichler and McIntyre, 1977; Allendorf and Phelps, 1980; Ryman and Ståhl, 1980; Verspoor, 1988; Busack and Currens, 1995; Campton, 1995; Norris *et al.*, 1999; Ford, 2002), but also marine species (Iguchi *et al.*, 1999; Sekino *et al.*, 2002).

Life history variables with an additive genetic variance (e.g. Reisenbichler and McIntyre, 1977; Cross and King, 1983; Taniguchi *et al.*, 1983; Hard, 1995), or developmental and morphological traits (Leary *et al.*, 1985) are also subject to change. Captive breeding and hatchery programmes also can lead to elevated frequencies of deleterious alleles that are otherwise kept at low frequencies in wild populations by selection (Lynch and O’Hely, 2001). Hybridization of genetically altered individuals with wild individuals can lower the fitness of offspring (outbreeding depression). Even if cultured individuals have not been genetically altered, supplementation may still pose a genetic threat to wild populations (Box 6).

### 5.4 Hybridization and outbreeding depression

Hybridizations between genetically divergent wild populations can occur for several reasons (Epifanio and Nielsen, 2001). One is the inadvertent or intentional introduction of genetically divergent conspecific individuals into a native population. Non-native individuals can be inadvertently introduced by ship ballast water, or as escapes from mariculture or aquaculture. Less common are natural or intentional habitat modifications that bring previously isolated populations in contact with one another. Ecological or competitive interactions between introduced and native individuals may drive wild populations to extinction. Genetic effects, although less obvious, can be equally detrimental to the survival of a species or stock. Genetic changes are greatest in captive populations closed to wild individuals. These results indicated that releases of
Ryman and Laikre (1991) outlined how stock supplementations can reduce genetic diversity through “genetic swamping”, even though the census size of the population in the wild increases. Captive individuals are generally produced from only a small number of parents relative to the number of potential parents in the wild. Releases of cultured individuals increase the parent-offspring variance and reduce the effective population size of the wild population, even though census numbers may be larger. Ryman and Laikre (1991) found that the effective population size equaled the sum of wild (Nw) and captive (Nc) parents only when the fraction of captive progeny was Nc/(Nc + Nw). Effective population sizes at other values of Nc and Nw are smaller. Importantly, supportive breeding in most instances reduces the total effective population size below what it would have been without supplementation. These smaller effective population sizes can lead to a loss of genetic diversity. Genetic swamping is a concern in species with high fecundities and high larval or juvenile mortality rates, a characteristic of most marine species.

Waples and Do (1994) explored this effect in more detail for Pacific salmon. They found that the extent of genetic swamping depended on the number of parents used in culture and not on the fraction of the wild population used for spawning. Genetic swamping can be hastened when individuals of hatchery origin are included a broodstock (Figures C and E). The most important determinant of levels of inbreeding in wild populations is the size of a wild population after supplementation. Even if the supplementation is successful and the wild population remains large, continued supplementation will eventually lead to the complete replacement of wild individuals with hatchery descendents. One of the few attempts to monitor the effects of supplementation showed reductions in genetic diversity in some populations of brown trout (Salmo trutta) that are likely due to stocking (Hansen et al., 2000).

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**BOX 6**

**Genetic effects of supplementation**

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Figure explanations (A-E):
captive individuals can pose a genetic risk to wild populations through hybridization (Box 7).

In a survey of the literature on hybridizations in fish, Utter (2001) found several generalizations (modified here).

1. Freshwater species are more susceptible than anadromous species to introgression from distinct lineages. Anadromous species are adapted to a greater number of life history variables (freshwater migration timing, marine migration, natal homing, run timing) than are freshwater species. Hence, introgression may be prevented by outbreeding penalties against hybrids in anadromous species.

2. Genetic distances between lineages of freshwater fishes are poor predictors of introgressive hybridization. Hybrids have occurred between highly diverged lineages and even between species.

3. Anadromous populations may be more prone to displacement than to introgression between major lineages. However, introgression commonly occurs between subgroups within major lineages.

4. Persistent disruption of subgroup adaptation through hybridization with non-native individuals retards the full potential for productivity of natural populations.

When selective pressures on captive populations are not managed, or when introgressive hybridizations are recurrent from long-term supplementation releases, genetic transformations of wild populations can potentially lead to the inability of a wild population to sustain itself without supplementation (Lynch and O’Hely, 2001).
BOX 7
Genetic effects of hybridization

The major genetic risk of hybridization is the disruption of adapted gene complexes and loss of fitness (outbreeding depression) (Rhymer and Simberloff, 1996). In one form of outbreeding depression, native individuals are better adapted to particular habitat conditions than are either the introduced or hybrid individuals. For example, experimental hybrids between even- and odd-year run pink salmon (Oncorhynchus gorbuscha) showed much lower survival rates than either of the two control groups (Gharrett and Smoker, 1991). Outbreeding depression can also occur in hybrids between geographically separated groups of the same year type (Gilk et al., 2004). A second form of outbreeding depression occurs when non-native genes are introduced into the genomes of wild individuals after the first generation of hybridization (introgression). Introgression disrupts the genes influencing a particular adaptation. Depending on the mode of expression of the genes, first generation hybrids may not be affected, but genetic recombination during reproduction separates co-adapted genes on parental chromosomes and reduces fitness in the introgressed individuals.

Reduced hybrid fitness has been documented experimentally in ‘common garden’ experiments for numerous freshwater (e.g. Dowling and Moore, 1985; Philipp et al., 2002; Neff, 2004) and anadromous fishes (e.g. Ferguson, 1986; Hawkins and Foote, 1998; Leary et al., 1985; McGinnity et al., 2003). A much longer list of species shows evidence of introgression from molecular markers (see Utter, 2001). However, virtually no examples exist of outbreeding depression in marine fishes, even though hybridizations are well documented with molecular methods. One reason for the apparent lack of outbreeding depression in marine fishes may be that it is difficult to demonstrate outbreeding experimentally. Another reason may be that local adaptations are not as prevalent in marine species because high levels of gene flow may prevent local adaptations. Generally, the lower levels of genetic divergence between populations of marine fishes and many invertebrates as detected by molecular genetic methods indicate high levels of gene flow. In marine species, substantial supplementation efforts have not always resulted in the expected increases in population abundance (Larkin, 1991; Masuda and Tsukamoto, 1998). These failures could in part be due to undocumented introgressive hybridization with long-term releases of cultured individuals.

6. MAPPING THE POLICY ENVIRONMENT
Numerous national and international initiatives have been proposed to explore ways of reversing declining abundances of the world’s biological resources (Table 2). Chief among these is the 1992 Convention on Biological Diversity (CBD, 1993), which calls for the conservation of biological diversity at three levels: genetics, species and ecosystems. Kenchington et al. (2003) noted that most initiatives focus on the conservation of species and ecosystems with little attention to genetics. Although ecosystem-based fishery management promotes the preservation of ecosystems and represents a major step toward achieving sustainable uses of natural resources, ecosystem management may not always protect genetic diversity within a species.

In addition to international initiatives, many countries or national organizations have outlined specific national problems and have attempted to implement policies intended to protect biodiversity at several levels. Parts of these policies have been formulated to address genetic issues directly. For example, the Fisheries Society of the British Isles recently published a briefing paper (FSBI, 2004) outlining the effects
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<th>Programme or Declaration</th>
<th>General intent</th>
<th>Statements or implications for genetics</th>
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<tbody>
<tr>
<td>1993 FAO Compliance Agreement</td>
<td>Promote compliance with international conservation and management measures by fishing vessels on the high seas.</td>
<td>Indirect: Reduce fishing pressure on harvested species.</td>
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<td>1995 Kyoto Declaration</td>
<td>Underscore importance fisheries to food security in developing countries.</td>
<td>Indirect: Sustainable fisheries.</td>
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### TABLE 2 (Cont.)

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<th>Programme or Declaration</th>
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<tr>
<td>2001 Reykjavik Declaration on Responsible Fisheries in the Marine Ecosystem</td>
<td>Recognizes fisheries impact on ecosystem, and hence, on fishery productivity.</td>
<td>Indirect: Preservation of ecosystem services and integrity of fishery populations.</td>
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### Organizations

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<th>Programme or Declaration</th>
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<td>1945 FAO established as specialized agency within United Nations</td>
<td>Provide forum to address issues relating to development and sustainable use of living marine resources. Provide fishery databases to support formulation of fishery management policies.</td>
<td>Direct: Numerous publications on importance of genetic processes in management and sustainable use of marine resources.</td>
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<td>1948 World Conservation Union (IUCN). World's largest conservation network, bringing together 82 States, 111 government agencies, more than 800 non-governmental organizations (NGOs).</td>
<td>Influence, encourage and assist societies throughout the world to conserve the integrity and diversity of nature and to ensure that any use of natural resources is equitable and ecologically sustainable.</td>
<td>Direct: Genetic diversity is one of the three forms of biodiversity recognized by the World Conservation Union (IUCN) as deserving conservation.</td>
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<td>1983 FAO Commission on Genetic Resources for Food and Agriculture</td>
<td>Permanent forum where governments discuss and negotiate matters relevant to genetic resources for food and agriculture.</td>
<td>Direct: Discussion of policies and practices influencing genetic diversity in plant and animal resources.</td>
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### TABLE 2 (Cont.)

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<tr>
<td>World Resources Institute (WRI)</td>
<td>Promote sustainable use of living resources through dissemination of information.</td>
<td>Indirect: 1. Reverse ecosystem degradation. 2. Protect global climate system.</td>
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<tr>
<td>1995 UNEP Global Programme of Action (GPA)</td>
<td>Protection of marine habitats from land-based activities</td>
<td>Indirect: Ecosystem protection</td>
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<tr>
<td>1994 International Coral Reef Initiative (ICRI)</td>
<td>Protection and restoration of reef ecosystems.</td>
<td>Indirect: Recognition that reefs are important fish nursery areas.</td>
</tr>
<tr>
<td>Joint Group of Experts on the Scientific Aspects of Marine Environmental Protection (GESAMP)</td>
<td>Provides advice on impact of human activities on marine ecosystems.</td>
<td>Indirect: Population health through sustainable use.</td>
</tr>
<tr>
<td>Marine Protected Areas (MPAs) initiated by World Bank, World Conservation Unions (IUCN), Great Barrier Reef Marine Park Authority (GBRMPA) and Global Environmental Facility (GEF).</td>
<td>Establishment of marine protected areas to aid in habitat and species restorations.</td>
<td>Indirect: Restoration of populations through protected marine areas.</td>
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of fishing on biodiversity in the North Sea and highlight specific threats to genetic diversity. Elsewhere, endangered species legislation in the United States (Endangered Species Act) has been interpreted by government conservation agencies to protect genetic diversity within and among intraspecific population groups (Waples, 1991). Box 8 gives an example of the use of this legislation to extend protection to threatened population groups of Chinook salmon (*Oncorhynchus tshawytscha*) in western North America.

7. DEVELOPING A FRAMEWORK FOR CONSERVING GENETIC DIVERSITY

It is clear from the arguments presented here that preserving genetic diversity in natural populations subject to capture fisheries is important for maximizing harvests and achieving the sustainable use of fishery resources. Four steps can be taken to develop a framework for conserving genetic diversity in capture fisheries (modified from Kenchington *et al.*, 2003):

**Step 1: Identify management objectives**

The rationale for developing the objectives of a conservation or management programme differs among disciplines and determines the directions of management efforts. Goals can be evaluated by arguments from several broad disciplines representing perspectives from the past, present and future (Bowen and Roman, 2004). A systematist may argue that a major goal should be the conservation of species representing the heritage of past evolutionary diversifications (Forey *et al.*, 1994; Wheeler and Cracraft, 1996; Vecchione *et al.*, 2000; Bowen, 1999).

From a contemporary perspective, an ecologist might argue that preserving functional ecosystems is the best way to conserve the components of genetic diversity among species and among conspecific populations. Changes in one component of an ecosystem by overfishing, for example, can ripple through an entire system and threaten the stability of species not targeted by a fishery. The loss of an ecological component in an ecosystem can have often unpredicted effects on other parts of an ecosystem (Brodziak and Link, 2002). A sociologist might argue more narrowly for the preservation of genetically influenced traits in a species or population that is valued by society. An economist might argue for the preservation of specific genes with potential pharmaceutical or commercial value.

With an eye to the future, an evolutionary biologist might argue for preserving the breadth of genetic diversity in a species to ensure its capacity to adapt to future environmental changes (Crandall *et al.*, 2002; Bowen and Roman, 2004). The rationale many conservation efforts is the preservation of genetic diversity to allow future adaptive shifts (e.g. Waples, 1995). Ecological and evolutionary considerations, however, are views of the same events on different temporal scales (Frank and Leggett, 1994). All these arguments must be weighed openly by society to set conservation priorities and to provide a foundation for setting management objectives.

In practice, conservation and management goals are often forged by the contradictory demands of industry, politicians, economists, ecologists and conservationists. Unlike conservation efforts, which are often directed at preserving components of genetic diversity, the goals of managing large fisheries are not usually directed at preserving genetic diversity itself, but at the population processes influencing this diversity. An underlying objective might be to maintain populations in a natural setting that allows ‘normal’ ecological and evolutionary processes to occur and to maintain the full geographical range of a species (Thorpe *et al.*, 1995; Taylor and Dizon, 1999). Other management objectives might include an increase in recruitment or a reversal of the effects of selective fishing on average size, maturation age or spawning timing.
Run timing: closed circles, spring; open square, summer; open circle, fall; asterisk, winter. Twelve geographic provinces (A–L) were delineated with allozyme frequencies and life-history information, such as spawning migration timing and the length of juvenile freshwater residence (Waples et al., 2004).

The “Endangered Species Act” (ESA) of 1973 in the United States of America mandated that endangered or threatened species be identified for special conservation efforts. Waples (1991) developed a framework to identify “distinct population segments”, which could be considered to be “species” under the ESA and receive the same protections as an endangered species. This framework invoked two criteria based on genetic and evolutionary considerations. A population represented a distinct population segment if it was reproductively isolated from other populations in the same species and if it represented an important component of the evolutionary legacy of a species. Genetic, ecological, geographical and life-history information was used to evaluate the statuses of populations in seven species of anadromous salmonids inhabiting western of the United States of America. While coastal fisheries can potentially limit the abundances of salmon populations, spawning biology and early life-history stages appear also to be important limiting factors.

**Step 2: Assess genetic risk**

This step is related to the first step. A clear understanding of the risks associated with the loss of genetic variability through inbreeding and stock extirpations, or the disruption of genetic structure through hybridizations, will help to guide the development of management goals. One important research agenda addresses the extent and rapidity of adaptation in local populations and the extent that human activities disrupt local adaptation (e.g., Taylor, 1991; Miller and Kapuscinski, 1994; Currens and Busack, 1995;
Sheridan, 1995; Conover, 1998; Law, 2000). Answers to these questions bear on the extent that evolutionary processes should be factored into management objectives. A growing body of evidence indicates that fish are often finely adapted to local habitats (Gilk et al., 2004; Utter, 2004), and that genetic changes can occur rapidly after transplantation (e.g. Kinnison et al., 1998), in culture (e.g. Hindar et al., 1991) or in response to fishing selectivity (e.g. Heino, 1998; Stokes and Law, 2000). Other genetic risks may come from intraspecific hybridizations between wild and fish that have been genetically modified in captivity (e.g. Leary et al., 1985; Philipp et al., 2002).

**Step 3: Identify reference points**

Setting benchmarks to evaluate progress toward fulfilling management objectives is a critical step in the process. Reference points have been defined by an ICES working group (ICES, 2001) as “specific values of measurable properties of systems (biological, social, or economic) used as benchmarks for management and scientific advice”. The purpose of setting benchmarks is to increase the awareness of the consequences of inaction on a particular problem. Two kinds of reference points can be distinguished (ICES, 2001). The first are ‘target reference points’, which are properties of stocks, species or ecosystems that help to achieve biological, social and economic goals. The second are ‘limit reference points’, which are threshold values of resource variables that trigger a conservation concern of unacceptable risk or irreversible harm. The setting of reference points for the preservation of genetic diversity depends on defining particular genetic risks to short-term goals such as maintaining stock abundance, economic return and species survival, and to long-term goals of preserving the capacity to adapt to environmental change.

The challenge in setting reference points is to understand what facets of genetic diversity are important for achieving particular goals. Only an integrated research agenda that includes genetics, ecology and economics can provide this understanding. Even a basic understanding of some genetic mechanisms is lacking. For example, little is known about what levels of genetic diversity are needed for a species to thrive and adapt. Generally, the wisdom is that as much gene diversity should be conserved as possible and that the loss of diversity leads to reductions in production. Theoretical considerations indicate that populations should not drop below 1 000-5 000 individuals to minimize the loss of gene diversity through random drift (Lynch and Lande, 1998). Yet, counter examples show that species can thrive after experiencing bottlenecks in population size that eliminated nearly all genetic diversity (Hoelzel, 1999).

**Step 4: Monitor progress**

Patterns of genetic variability within and among populations of a species can be monitored directly with molecular genetic methods or indirectly with models and population baseline data. Surveys of molecular genetic variability are costly, but have been vital for estimating levels of connectivity among and gene diversity within populations. Temporal sampling is needed to monitor the effects of management actions, after an initial survey establishes a baseline. One constraint on genetic monitoring is the lack of historical data. Major declines in stock abundances from fishing occurred several decades before molecular methods were first used to survey genetic variability on a large scale in the 1970s. Hence, a pre-fishing baseline is difficult to establish for most species. Temporal datasets have been instrumental in showing gene-frequency shifts in some species that appear to be due to fishing intensity (Hauser et al., 2002). Another constraint in using molecular genetic markers to monitor the effectiveness of management on short time scales is that genetic profiles may not respond rapidly to environmental and demographic events. The development of high throughput methods of surveying genetic diversity (e.g., SNPs, Smith et al., 2005; DNA microarrays, Cossins and Crawford, 2005) will make genetic monitoring more feasible in the future.
Avise (2001) outlines how cyto-nuclear signatures of genetic variability can be used to detected hybridizations and introgressions. Dowling et al., 2005 provide an example of monitoring the genetic effects of supplementing over 11 years populations of endangered Catostomid fish.

Genetic parameters can also be monitored indirectly with theoretical models and data for population abundance, population demography and geographical distribution. For example, population models indicate that strong reductions in population size, metapopulations extinction dynamics and population fragmentation can lead to the loss of genetic variability.

8. CONCLUSIONS

The relative importance of genetic processes in species supporting capture fisheries differs among ecosystems and species. Fisheries in rivers and lakes are largely focused on species with naturally fragmented populations. Dams and land transformations further isolate some populations by destroying migration corridors between populations. Inland species are therefore vulnerable to the loss of genetic diversity through the metapopulation processes of extinction and colonization and through random drift in small populations. Hence, genetic concerns are largely addressed under the framework of conservation biology and theory relating to inbreeding and inadvertent hybridization.

Anadromous species support large commercial fisheries in the North Atlantic and North Atlantic. These species are especially vulnerable to ecological disturbances because of their complex life-history cycle, which spans freshwater and marine habitats. The loss of between-population genetic diversity through population extinctions in some species is especially acute in areas of extensive human development and degraded habitats. Native salmonids appear to be particularly at risk from hybridizations with genetically divergent conspecific individuals.

In the marine realm, species supporting capture fisheries also face genetic threats. Marine species in open waters can experience the loss of genetic diversity through the extinction of local subpopulations, genetic swamping through stock enhancements with artificially propagated individuals and intentional or inadvertent introductions of related species. The loss of genetic variability through random genetic drift appears to be less important for marine species than for inland and anadromous species. Even so, shifts in gene frequencies from drift induced by overfishing have been documented in some marine species and imply reductions to small effective population sizes.

Overfishing, habitat degradation and climate shifts appear to be far more important threats to stocks of most species than are genetic risks. This assessment is illustrated by a recent collection of papers from a symposium, entitled "Fisheries, past, present and future" (Philosophical Transactions of the Royal Society B, 2005, vol. 360), which did not include a single article dedicated to genetic issues in the management of capture fisheries. Although genetic processes were briefly discussed in some of the 14 major articles, highlights of genetic issues were absent from the introductory summary of the conference (Beddington and Kirkwood, 2005). Genetic processes appear to be perceived as far less important than ecological and life history processes influencing stock abundances.

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ANNEX 1

METHODS AND CONCEPTS FOR SURVEYING GENETIC RESOURCES

1. GENETIC BASIS FOR MOLECULAR VARIABILITY
The genetic information needed for the development and physiological maintenance of an individual is stored in a long polymeric molecule called DNA. DNA is found in two organelles in a cell: over 99% of DNA is located in the nucleus, but a small fraction occurs as a plasmid-like circular structure in mitochondria (Figure a). Genes encoded by nuclear DNA are inherited from both parents, and hence occur in pairs to form a diploid genotype (Figure b). Genes encoded by mitochondrial (mt) DNA, however, are maternally inherited in most species and hence occur as a single haplotype in an individual. The analysis of mtDNA, which also lacks recombination, can provide unique insights into population structure that is not possible with nuclear DNA (Avise, 1994). The entire complement of DNA is denoted by the term genome, and various parts of the genome serve different functions.

One important function is to encode information that can be translated into proteins. The coding parts (exons) of many genes are often interspersed by noncoding (introns) sections of DNA. Introns are less constrained by natural selection and hence mutate at a higher rate than the protein coding portions of a gene. Other parts of the genome encode regulatory information, important in development and gene expression. A large portion of the genome appears to serve no coding function, but may be important in the physical arrangement of DNA in the nucleus. These sections of DNA often have large numbers of short repeats called microsatellites.
2. TRENDS IN THE DEVELOPMENT OF MOLECULAR GENETIC MARKERS

Early methods of surveying genetic variability, such as immunological assays and allozyme electrophoresis, examined the products of DNA coding genes. A large amount of information on the genetics of natural and cultured populations of aquatic organisms has been produced since the early 1970s, when protein electrophoresis was first used on a large scale to the survey genetic variability in and among natural populations. However, technological developments since the 1980s have produced methods that assay DNA polymorphisms directly (Palsbøll, 1999). The most important population markers include direct sequencing, restriction enzyme fragment length polymorphisms (RFLPs), amplified fragment length polymorphisms (AFLPs) and single nucleotide polymorphisms (SNPs). The application of these methods has been greatly aided by the development of the polymerase chain reaction (PCR), which amplifies targeted DNA sequences from small amounts of tissue. These various techniques provide complementary information about natural populations and are variously suited to answering different questions. When molecular population markers are under natural selection, caution must be used to infer movement between subpopulations from the geographical distributions of allele or haplotype-frequencies. Although selection on allozymes can occur because they encode proteins (Mitton, 1997), DNA cannot always be assumed to be free of selection (Avise, 1994; Bazin et al., 2006; Nielsen et al., 2006). Biogeographical and laboratory evidence indicates that selection may shape the geographical distributions of protein (Powers et al., 1991; Powers and Schulte, 1998), mitochondrial DNA (Árnason, 2004; Bazin et al., 2006; Grant et al., 2006) and nuclear DNA variants (Pogson and Mesa, 2004; Canino and Bentzen, 2004; Case et al., 2005). The nature and intensity of selection must be understood when using selected population markers to infer population structure. Patterns of divergence for adaptive and neutral markers may not coincide (McKay and Latta, 2002). Although selectively neutral molecular markers will continue to be important, Vrijenhoek (1998) argues that adaptive traits should also be examined to help resolve conservation and management problems.

3. ESTIMATING EFFECTIVE POPULATION SIZE

One promising use of genetic data is to estimate the effective sizes of fishery populations. The genetic concept of effective population size is the number of individuals that actually contribute genetic information to the next generation. Not all individuals in a population produce offspring that reach reproductive maturity. In the marine environment, many species show a large variance in family size because of variability in the physical and biological factors influencing larval survival (Hedgecock, 1994). Fishery resource managers, on the other hand, focus on the actual number of individuals in a population (census size). The difference between these two numbers can be large for the same stock. Census population sizes are generally at least ten times the effective sizes (Frankham, 1995; Nunnery and Elam, 1994). In some cases, census size can be as much as three orders of magnitude larger than effective population size (Turner et al., 2002). The loss of genetic diversity has been detected in some species, even though census numbers may still be large (Hauser et al., 2002). Effective population size can be estimated in several ways. One way is to estimate the drift effective size by examining temporal changes in gene frequencies. The concept behind this approach is that effective population size influences the amount of genetic drift in a population. Small populations experience a greater amount of genetic drift, and hence greater gene frequency changes, than do larger populations. This method requires gene frequencies estimates from different generations. For some species, the analysis of archived fish scales from collections in the last few decades has provided estimates of historical gene frequencies (Miller and Kapuscinski,
Different statistical approaches have been used to extract unbiased estimates of effective population sizes from gene frequency data (Luikart et al., 1999; Wang, 2001; Berthier et al., 2002). These methods provide estimates of effective population sizes of contemporary or very recent populations. Other methods of estimating effective population size use equations from evolutionary theory that incorporate long-term effective population size. One approach is to use observed heterozygosity, which is expected to be a function of effective population size and the neutral mutation rate (Waples, 1991). Another approach is to estimate the coalescence times for mtDNA haplotypes when recombination is absent (Avise et al., 1988). Coalescence time (the time until haplotype lineages trace to a common ancestral haplotype) is expected to be a function of population size. While these estimates may be reveal long-term features of the population biology of a species, they are not always useful for making management decisions, because they may not represent current population sizes.

4. REFERENCES


ANNEX 2

1. COMPUTER PROGRAMMES FOR GENETIC ANALYSIS

The development of new technologies to detect molecular variation and automation of several steps in these laboratory analyses have led to the production of large amounts of genetic data. The availability of these data has stimulated the development of new statistics and computer programmes, which provide insights from data not previously possible (Zhang and Hewitt, 2003). The use of computers provides the opportunity to test hypothesis with bootstrapping and coalescent simulations, in addition to standard parametric, non parametric and exact tests.

Most, if not all, the computer programmes available for genetic analysis can be downloaded from the web sites of academic institutions without charge. Several groups of programmes are available. Multipurpose programmes are generally used to examine genotype or sequence data and to describe gene diversities within and among samples. These programs include ARLEQUIN (Excoffier et al., 2005), DnaSP (Rozas et al., 2003), FSTAT (Goudet, 1995); GENEPOP (Raymond and Rousset, 1995), GENETIX (in French only; Belkhir et al., 2000), and MEGA (Kumar et al., 2004), among others. The basic facilities offered in these programs are reviewed in Excoffier and Heckel (2006).

In addition to these basic programmes, many other programmes incorporate algorithms that attempt to assign individuals to particular populations. These include BAPS (Corander et al., 2004), GeneClass (Piry et al., 2004) and GeneLand (Guillot et al., 2005), STRUCTURE (Pritchard et al., 2000), among others. Also in this group are programs written for fishery management to estimate the origins of individuals in areas of stock mixing (BAYES, Pella and Masuda, 2001; WHICHRUN, Banks and Eichert, 2000). Hansen et al., (2001) reviews the utilities of these and other mixed-stock computer programmes for microsatellite DNA markers.

Another group includes specialized programme performing a variety of tests of past demographies (ARLEQUIN; DnaSP; BATWING, Wilson et al., 2003, among others). Algorithms in these programmes search for evidence of population growth or bottlenecks in population size. Molecular markers are often assumed to be neutral to the effects of selection. This assumption can be tested by gene- or haplotype-frequency distribtuions (MEGA; FDIST2, Beaumont and Nichols, 1996). Migration is also an important factor shaping the genetic population structure of a species. Estimates of migration between populations (gene flow) are often used in devising conservation and management strategies (COLONISE, Foll and Gaggiotti, 2005; MIGRATE, Beerli, 2006, among others).

Inferring phylogenetic relationships among species can also be important to the management of a multispecies fishery. Phylogenetic trees were first constructed from genetic distances estimated from gene frequencies. The widespread availability of DNA sequences, however, allow more sophisticated approaches to tree construction (see Felsenstein, 2003). These methods include parsimony, maximum likelihood and Baysian algorithms (Nei & Kumar, 2000). Many of the general programmes listed above provide options to use some of these methods. However, several specialized programmes can be downloaded from the internet. Some of the more commonly used programmes include PHYLIP (J. Felsenstein: http://evolution.gs.washington.edu/phylip.html), PAUP® (D. Swofford: Sinauer Associates), MacClade (W. Maddison: http://phylogeny.arizona.edu/macclade/macclade.html) and DAMBE (X. Xia: http://aix1.uottawa.ca/~xxia/software/software.htm), among others.

Many situations encountered by fishery biologists do not easily fit the assumptions of some biological and statistical models, which are often simplified for easier use. For example, equal population sizes and equal migration rates between populations are assumed in several genetic population models, but in nature are seldom equal to one
A refinement in the use of statistical models to interpret genetic data is the use of simulation programs to model particular situations (e.g., SIMCOAL, Excoffier et al., 2000; EASYPOP, Balloux, 2001; METASIM, Strand, 2002; MESQUITE, Madison and Madison, 2004; among others). Future approaches to DNA data analysis will use maximum likelihood and Bayesian methods tailored to particular situations (e.g., Whitlock and McCauley, 1999; Pritchard et al., 2003; Dawson and Belkhir, 2001; Wilson and Rannala, 2003).

These computer programmes should be used cautiously. The use of some programmes is complicated by the different input formats. This requires reformatting of datasets manually or with programmes designed for data conversion (see Excoffier and Hackel, 2006). Another caution is that different programmes may produced different values of the same statistics for the same set of data. This is likely due to differences in how the programmes are written. Lastly, these programmes offer numerous options for analyzing data and produce a wealth of statistical output. A researcher should always take the time to read the background literature on how a statistic is calculated and its interpretation. User documentation of some programmes (e.g. ARLEQUIN) presents some explanations. However, the successful application of many programmes requires that the user read the original literature.

2. REFERENCES


ANNEX 3

USE OF GENETIC DATA IN FISHERY MANAGEMENT

1. DEFINING POPULATIONS FOR MANAGEMENT
One problematic issue has been a lack of consensus on the definition of a population, even though the ‘population’ is a fundamental unit in ecology, evolution and fishery management. In fishery management, few definitions of a population are operational enough to be used objectively by researchers or policy makers (Waples and Gaggiotti, 2006). Yet, how populations are connected to one another through migration has important consequences for devising management plans. Fishery managers usually agree that management units should coincide with natural population partitions, but how natural populations are defined is a subject of continuing debate (Ryder, 1986; Moritz, 1994; Waples and Gaggiotti, 2006; Schaefer, 2006; Palsbøll et al., 2006).

Two contrasting, but overlapping, views appear in conservation biology and fishery management. In one perspective, the conservation of populations representing major evolutionary lineages is thought to be important. Beyond the conservation of evolutionary legacy is the attempt to maintain the population processes that produce deep levels of diversity in a species (Moritz, 2002). Genetic variability has to be conserved to allow a species to adapt to environmental changes taking place on decadal and millennial time scales. On the other hand, the needs of fishery management are short term and require a greater resolution of population structure on smaller geographical and temporal scales. To this end, traits responding rapidly to environmental variability such as morphology, meristic counts and life history patterns have frequently been used most to define populations and stocks.

A major problem arises in the use of genetic methods for conservation and fisheries resource management, because evolutionary and ecological definitions of a population are mistakenly used interchangeably (Figure 4). Both kinds of populations are defined by the degree of connectivity among populations through the exchange of migrants (population structure). However, far less migration is required to maintain genetic cohesiveness among populations on evolutionary time scales than is required to produce demographic homogeneity among populations. The evolutionary population concept predominates in conservation biology, in which a major concern is the protection of genetic lineages, which allows a species to adapt to environmental changes. In fishery management, an ecological population may not be genetically distinctive, but may still show life history differences or geographical isolation requiring separate management.

Ecological definitions of a population, however, focus on interactions between individuals that influence the demographic characteristics of a population, including competition, age structure and birth and death rates. These kinds of variables are used to define stocks for the biomass assessments used to set harvest limits. From this ecological viewpoint, demographic independence between populations can still persist with much large amounts of immigration (Figure 4). A limited amount of data indicates that demographic independence between populations occurs when the proportion of immigrants (m) falls below 10% (Hastings, 1993). The key variable for management considerations, especially of marine species, is the proportion of migrants, m. However, models used to interpret genetic data yield only estimates of Nm, the number of migrants between populations. Another difficulty is that N represents population size integrated over recent evolutionary time and not necessarily the size of a contemporary population. Estimates of census size are also not useful for estimating N, because effective population sizes may be an order of magnitude smaller than census size (see below). Future research will focus on the development of simulation and modelling tools that integrate ecological and genetic data for particular situations.
2. MODELS OF GENETIC POPULATION STRUCTURE
The origins of genetic population structure in freshwater and riverine organisms are fairly well known. Both demographic and genetic populations are usually delimited by lake shorelines and watercourses that represent strong physical barriers to migration. Unexpected genetic similarities between populations can usually be explained by historical events, such as headwater captures, altered river drainages or gene flow in proglacial lakes after the last ice age. Some of the classic models of population structure [e.g. island model of migration (Wright 1940)] have been used effectively to estimate contemporary levels of migration in many species from molecular markers (Neigel, 1997). Populations of marine fishes, on the other hand, are not generally bounded by strong barriers to migration. Nevertheless, the genetic structures of marine populations can be influenced by spawning migrations (Ruzzante et al., 2006), sea surface temperature gradients (Beare et al., 1995; Attrill and Power, 2002) and currents and ocean frontal systems (Rocha-Olivares and Vetter, 1999; Shaw et al., 2004).

Two contrasting models of population structure have been posed for marine fishes. Under a local, self-sustaining model (vagrant-member hypothesis, Sinclair, 1988), populations are adapted to local environmental settings and persist only when conditions allow uninterrupted cycles of spawning, growth and reproduction. The magnitude of gene differences between populations depends on effective population size, which influences genetic drift and gene flow, which tends to reduce population differences. An extension of this model incorporates the ephemeral nature of some populations by recognizing that regional extinctions and colonization occur in marine populations (Smith and Jamieson, 1986; McQuinn, 1997). This model predicts that subpopulation extinctions and recolonizations should lead to a mosaic pattern of genetic diversity and gene frequency heterogeneity.

Under the ‘basin model’ (MacCall, 1990), the most favourable habitats lie toward the centre of a species’ geographical range. During ocean-climate instability, subpopulations at the edges of the species’ range disappear by contracting toward the central favourable basin. Abundances in the centre of the range remain constant, even though the species as a whole is declining. This model is also echoed in the classical biogeographical models of Mayr (1970), who found that environmental factors were most important in regulating abundances at the periphery of a species’ range. When these species represent ‘straddling stocks’, international cooperation is especially important in setting harvest limits. Subpopulations following this model might show gradients in gene diversity with the highest diversities in the centre of the range, where populations have been most stable. Considerations of these two models are important to formulating management policies and planning locations of marine protected areas.

Genetic data generally fail to show the high degree of isolation in open-ocean marine fishes predicted by Sinclair (1988). Genetic estimates of gene flow are high in most marine species (Table 1), implying the movements of tens and hundreds of individuals between subpopulations. Mitochondrial DNA data appear to support the basin model for California anchovy (Lecomte et al., 2004), but support a mosaic model for European anchovy (Grant, 2005; Magoulas et al., 2006). However, finer-scale differences have been detected among populations that are not isolated by obvious physical or hydrographic barriers (Hedgecock et al., 1994; Ruzzante et al., 1999). This chaotic variability is likely due to large reproductive variances among families (Hedgecock, 1994), rather than to isolation or adaptations to particular open-water habitats. The instability of marine waters on annual, decadal and millennial time scales likely prevents adaptations to specific areas. On a decadal scale, anchovy populations, for example, respond rapidly to small climate changes with range contractions and expansions (e.g. Cushing, 1982; Beare et al., 2004).
3. LOSS OF GENETIC DIVERSITY

A major concern in conservation biology is the maintenance of genetic diversity (measured by average heterozygosity). Values of heterozygosity, however, are affected most by the frequencies of abundant genes in the general range of 0.10–0.90. Genes at low frequencies contribute little to heterozygosity. Ryman et al. (1994, 1995) pointed out that the store of gene variability represented by low-frequency genes may be important in adapting to changing environments. Large populations have a greater capacity for retaining low-frequency genes than small populations, but the relative loss of rare genes during a population crash is much greater in large populations. For example, consider two populations of sizes 10,000 and 100,000,000, which are at mutation-drift equilibrium, but which are reduced to 1% of their original size (1,000 and 1,000,000). The loss of heterozygosity is negligible in both populations, but the small population retains 98% of the original gene number, while the large population retains only 1% of its genes.

4. REFERENCES


1. SUMMARY

The deep sea is the largest habitat on earth, covering around 53% of the sea’s surface, from the poles to the tropics. The deep-sea region starts at the shelf break at the continental margins, around 200 m, and extends down the continental slope and the continental rise to the abyssal plain at around 6 000 m, and the deep trenches. Deepwater fisheries occur on the continental slopes and on seamounts and exploit resources down to ~2 000 m. The continental slopes cover about 8.8% of the world’s surface, an area greater than all the continental shelves and shallow seas, and include the most variable habitats in the deep-sea with canyons, ridges, seamounts, hydrothermal vents, and cold seeps.

Definitions of deepwater fisheries vary geographically, but generally occur at depths greater than 400-500 m; trawl fisheries for orange roughy (Hoplostethus atlanticus) and ores (Pseudocyttus maculatus, Allocyttus niger and Neocyttus rhomboidalis) occur between 600-1 800 m, while long-line fisheries for toothfish (Dissostichus spp.) in the Southern Ocean operate down to ~1 800 m. Landings of deepwater fishes have risen from <0.5 m tonnes a year in the 1960s to >3  m tonnes by the late 1990s, with more than half of the annual catch taken from the Atlantic Ocean, but account for only ~5% of the total fish catch. The landing statistics are likely to be under estimates due to illegal, unreported and unregulated (IUU) fishing operations, and discards of bycatch species. Several deepwater fisheries have been characterized by “boom and bust” cycles. Catches of the armourhead (Pseudopentaceros wheeleri) on the North Hawaiian Ridge were estimated to have exceeded 150 000 tonnes a year during the late 1960s to 1970s where today no fishery exists. During the late 1990s a new fishery developed for orange roughy and alfonsino (Beryx spp.) in the South Indian Ocean with annual landings rising from <1 000 tonnes, peaking at 39 400 tonnes in 2000, and declining to <5 000 tonnes by 2002. In other regions orange roughy fisheries have been closed to commercial fishing, following a cycle of rapidly rising and declining catches. High catches of orange roughy in some areas have been maintained, at least temporarily, through local scale serial depletion as neighbouring seamounts and hills are fished down.

Deep-sea fishes include a large number of diverse species. Not all deepwater fishes are well described and molecular tools are being used to resolve taxonomic questions of species identity. Species exploited by deepwater fisheries include both shelf species, that extend down the continental slopes, and species restricted to depths >400-500 m. Most species are caught by trawls on seamounts and ridges, although line fishing and gillnets, and traps for invertebrates are used; toothfish (Dissostichus spp.) in the Southern Ocean are taken by trawl and long-line fisheries. An artisanal long-line fishery has existed for the black scabbard fish Aphanopus carbo for more than a century off Maderia, but most deepwater fisheries are relatively new and capital-intensive. A few small-scale
Deepwater fisheries occur where the shelf is narrow and the fishery areas are accessible by small vessels using drop lines. The sustainable yields from such fisheries maybe only a few hundred tonnes a year, but are important for small island states.

Deepwater fisheries generally target teleosts, with sharks taken as bycatch; only a few target invertebrates. In the North Atlantic deepwater fisheries, 22 species of teleosts 10 species of shark and two invertebrates (the red crab *Chaecon affinis* and the shrimp *Aristeomorpha foliacea*) make up the most important commercial species. Major species associated with seamounts include orange roughy, oreos, alfonsinos, and the roundnose grenadier (*Coryphaenoides rupestris*). A high degree of endemism has been reported for seamount invertebrates and fishes, but many of the targeted fish species have extensive ocean-wide and even cosmopolitan distributions.

As with coastal and shelf fisheries, conserving genetic diversity at the population, species, and ecosystem levels should be major goals for managing genetic resources in wild populations. Genetic issues identified for shelf species are likely to be magnified for deepwater species. Many slope and seamount species exhibit traits such as high longevity, slow growth rate, and late maturity, that make them more vulnerable to exploitation than most shelf species.

Marine fish tend to have higher levels of intraspecific genetic diversity than anadromous species, which in turn are more variable than freshwater species; a trend relating to larger evolutionary effective population sizes in marine fishes. Low levels of genetic diversity have been reported in the Antarctic toothfish *Dissostichus mawsoni*. Marine fishes show less spatial genetic differentiation than anadromous and freshwater species, due to the fewer barriers to gene flow in the marine environment.

A negative relationship reported between genetic differentiation and dispersal potential in coastal fishes appears to apply to deepwater fishes. Recent developments with new molecular tools, coupled with new analytical approaches, have revealed finer scale population structure within ocean basins for the Patagonian toothfish *D. eleginoides*, but for many deepwater fishes there is little or no information on genetic diversity within and among regions, and the scale of appropriate management units remain uncertain. Local declines among orange roughy fisheries on neighbouring seamounts suggest that they maybe independent units in the ecological time frame of fisheries management, in the absence of detectable genetic differentiation at small spatial scales.

Directional selection, through size-selective harvesting, has been implicated in changes in life history traits in heavily exploited stocks of shelf species, but has not been demonstrated in deepwater fishes, in part due to the limited time series of appropriate data. The genetic composition of a population can also change over generations due to random events. Changes due to genetic drift are most likely in small populations and are expected to be weak in marine fishes with large populations ($N > 10^6$). However 'sweepstake' events, due to high larval mortalities, can result in a small effective population size ($N_e$) several orders of magnitude smaller than the census population ($N$). Low $N_e / N$ ratios have been demonstrated in several shelf species and are equally likely to occur in some deepwater species, and potentially lead to loss of genetic diversity in collapsed stocks.

There is a general perception that the risk of extinction is low for commercially important marine fishes due to their large population sizes and wide geographical distributions. Only a few marine fishes have been listed as endangered and fewer appear to be close to extinction. Several traits of deepwater species (long life span, large body size, low natural mortality, and late sexual maturity) make them more vulnerable to extinction than shelf species, in particular those species that aggregate on seamounts. Deepwater fisheries have only been operating in the Northwest Atlantic Ocean since the 1970s, but already several species appear to meet the criteria of being critically endangered. Non-target species, that include teleosts endemic to
seamount complexes and elasmobranchs with low reproductive potentials, are also likely to be endangered.

Currently discarded fish waste from processing is used for low value products such as fish-oils, meals, pet foods, and silage. Bioactive compounds may be extracted from left-over fish-frames, internal organs, and invertebrate bycatch species for biotechnological and pharmaceutical applications, offering the opportunity to add value to fisheries. Some compounds derived from fish waste have been identified as potential nutraceuticals. Marine invertebrates that occur around hydrothermal vents may provide enzymes and biochemicals for the biotechnology industries and become target species in the future, raising further issues over exploitation of specialised deepwater habitats.

Genetic resources at the species and ecosystem levels are equivalent to ecological resources for which the management issues are well documented in the fisheries literature. The rapid development, and in some cases rapid depletion, of deepwater fisheries is of major concern to fisheries managers around the world, and has been identified repeatedly at local, regional, and international meetings. ICES have recognised that most exploited deepwater fishes are harvested unsustainably and radical reductions in fleets, in particular trawlers, are required to reduce effort and to conserve vulnerable habitats.

NGOs have expressed concern over the mortality of macro invertebrates taken as bycatch in deepwater trawl fisheries on seamounts, and for seabirds taken in toothfish trawl and long-line fisheries, although mitigation measures have been put in place to reduce the bird catch. The fragile and ancient coral “forests” found on seamounts that are amenable to trawling are quickly reduced to rubble by heavy trawl gear. Improvements to trawl gear and monitoring may allow the operation of deepwater pelagic trawls that avoid contact with bottom features. In the short term, one mitigation measure to protect vulnerable and unique habitats is to close selected areas to bottom trawling.

Many deepwater fisheries occur in high-seas areas compounding the problem of management and regulation. IUU fishing has been widespread in high seas fisheries. Increased surveillance and the introduction of a catch documentation scheme have reduced IUU fishing for toothfish within and outside the Commission for the Conservation of Antarctic Marine Living Resources (CCAMLR) Area in the Southern Ocean. Removal of fisheries subsidies should relieve pressure on deepwater stocks to some extent, but will need to be considered in parallel with other management tools. Regional Fishery Management Organisations (RFMOs) are being developed that cover high seas fisheries, and urgent action is required at the global level, to avoid shifting the deepwater fishing problem from one region to another. The inertia in developing and implementing international fisheries legislation, has lead to NGOs calling for the designation of large scale protected areas, and for a moratorium on bottom trawling. Recent initiatives have seen the establishment of a high seas benthic protected area in the Southern Indian Ocean, with further proposals for a network of large Marine Protected Areas or Benthic Protected Areas in waters around Australia and New Zealand, and in the Southern Ocean.

2. INTRODUCTION

The impacts of large scale industrial fishing on coastal ecosystems are well known. In some shelf and open ocean fisheries the community biomass has been reduced by up to 80 per cent within 15 years of exploitation (Myers and Worm, 2003). For the recorded species of coastal and oceanic sharks in the Northwest Atlantic Ocean fisheries, all except one have shown declines in relative abundance of >50% since the mid 1980s (Baum et al., 2003). Extinctions of marine species were thought to be rare events, but two species of skate are near extinction in the North Atlantic (Brander, 1981; Casey and Myers, 1998); in coastal fisheries the Pacific abalone *Haliotis sorensei* is in a
perilous condition (Davis et al., 1998), and some populations of giant clams, *Tridacna*, are locally extinct (Awaya and Lee, 2005).

Compared to the rest of the planet, much of the deepsea appears to be relatively unimpacted by human activities. International regulations prohibit the dumping of structures and radioactive waste in the deep-sea; and oil and gas extraction has been limited (Glover, AGS, C R 2003). However, the expansion of commercial fisheries into deep waters (>400 m) especially those in high seas outside coastal jurisdictions has been a major development in world fisheries in recent years (Watson, R and Morato, T 2004). Landings of deeper water species have increased, driven by technological developments, that enable fishers to target localised feeding and spawning aggregations of fishes in deepwater, and by catch-limits imposed in coastal fisheries. The biological issues of overfishing deepwater species are well documented and there is general agreement that many deepwater fishes are exploited beyond sustainable levels (Koslow, et al., 2000; Haedrich et al., 2001; Watson and Morato, 2004; FAO, 2005b; Devine, et al., 2006; Morato et al., 2006a and 2006b). Urgent action is required at the global level to reduce effort in deepwater fisheries and to protect fragile ecosystems from the impact of bottom trawling.

The impacts of fishing on deep-sea genetic resources are likely to be similar to those observed in shelf fisheries at the population, species, and ecosystem levels. In general, the effects of fishing on intra-specific genetic diversity are more difficult to detect than ecological impacts (Kenchington, E. 2003), but include loss of diversity through size-selective fishing favouring early maturity and slow growth (Dieckmann et al., 2006 in press), and through reduction in numbers of breeding fish (Hauser et al., 2002). Patterns of genetic diversity and population structure are not well known for most deepwater fishes. The life history traits of some deepwater fishes, in particular high longevity, slow growth, and late maturity, make them more vulnerable to fishing than coastal species (Morato et al., 2006a and 2006b), such that they may require different management systems (Clarke, M. et al., 2003). Some deepwater fishes already qualify as endangered species following <20 years of exploitation (Devine et al., 2006).

3. THE DEEP-SEA AND DEEPWATER FISHERIES

The deep-sea is the largest habitat on earth, covering around 53% of the sea’s surface from the poles to the tropics. The deep-sea region is generally recognised as starting at the shelf break at the continental margins (around 200 m) and extending down the continental slopes and the continental rises to the abyssal plains at around 6 000 m, to include the trenches. Much of the continental slope and the abyssal plain regions are covered by soft mud substrates, but the slopes include the most variable habitats in the deep-sea with canyons, ridges, seamounts, hydrothermal vents, and cold seeps. Four depth zones are recognised in the deep-sea: mesopelagic (200-1 000 m); bathypelagic (1 000-4 000m); abyssopelagic (4 000-6 000 m); and the hadalpelagic below 6000 m depth, in the deep ocean trenches.

The deep-sea is a dark, cold environment. There is no primary productivity via photosynthesis; even at depths of 150m light levels are just 1% of those at the surface and are insufficient to support photosynthesis. Concentrations of organic material decrease exponentially with depth, but the deep-sea is fuelled by a rain of sinking dead phytoplankton and nekton, and by many species that perform extensive vertical migrations, transferring surface production into deeper waters. Specialist habitats are maintained by organic material derived from sinking wood and from whale carcasses and have lead to the evolution of unique fauna (Distel et al., 2000) which along with fauna on hydrothermal vents and cold seeps are dependent on chemoautotrophic primary production (VanDover, 2000).
Issues, status and trends in deep-sea fishery genetic resources

Limited resources in shelf fisheries lead to an expansion of fishing effort into deeper waters during the 1980s. Some shelf fisheries expanded into deeper waters on the continental slope as technologies developed, e.g. the North Atlantic fishery for Greenland halibut *Reinhardtius hippoglossoides*. Other deepwater fisheries are relatively new and target species that are restricted to deepwater, e.g. the orange roughy *Hoplostethus atlanticus* found in the Atlantic, Indian and Pacific Oceans between 600-1 600 m.

Deepwater fisheries occur on the continental slope and on seamounts (see Box 1) and exploit resources down to ~2 000 m. The continental slopes cover about 8.8% of the world’s surface, an area greater than all the continental shelf and shallow seas (~7.5% of the world’s surface). Definitions of deepwater fisheries vary geographically; ICES uses the term deepwater fisheries for those in depths >400m; others define deepwater fisheries as those occurring deeper than ~500 m (Koslow *et al.*, 2000). Around New Zealand deepwater trawl fisheries occur between 750-1 500 m, while middle depths fisheries are recognised between 200-750 m.

Most deepwater species are caught by trawls on seamounts and ridges, although line fishing and gillnets are used, as are traps for invertebrates; toothfish *Dissostichus* spp. in the Southern Ocean are taken by trawl and long-line fisheries. An artisanal long-line fishery has existed for the black scabbard fish *Aphanopus carbo* for more than a century off Maderia, but most deepwater fisheries are relatively new, technology-dependent and capital-intensive. The Spanish deepwater fisheries are opportunistic with target species changing according to availability of other commercial species (Pineiro *et al.*, 2001). A few small-scale deepwater fisheries occur where the shelf is narrow and the fishery areas are accessible by small vessels using drop lines. The sustainable yields from such fisheries maybe only a few hundred tonnes a year, but are important for small island states.

Several deepwater fisheries have been characterized by “boom and bust” cycles. Catches of the armourhead *Pseudopentaceros wheeleri* on the North Hawaiian Ridge were estimated to have exceeded 150 000 t a year during the late 1960s to 1970s (Boehlert, 1986; Boehlert and Sasaki, 1988; Somerton and Kikkawa, 1992): today no fishery exists. Catches of the Pacific Ocean perch (*Sebastes alutus*) peaked at around 450 000 tonnes in the mid 1960s and have since fluctuated at 5-30 000 tonnes a year (Ianelli and Zimmerman, 1998). During the late 1990s a new fishery developed for orange roughy and alfonsino (*Beryx* spp.) on the Southwest Indian Ridge with annual landings rising from <1 000 tonnes, peaking at 39 400 tonnes in 2000, and declining to <5 000 tonnes by 2002 (FAO, 2002). In other regions orange roughy fisheries have been closed to commercial fishing, following a cycle of rapidly rising and declining catches. High catches of orange roughy in some areas are maintained, at least temporarily, through local scale serial depletion as neighbouring seamounts and hills are fished down.

The top fish species by landings (>1 m tonnes a year) that account for 30% of the total world capture fisheries are shelf and pelagic species, and of these 7 are fully exploited or overexploited (FAO 2005c). Furthermore, the top 71 species of fish and invertebrates, which account for ~50% of capture production by tonnage are shelf and pelagic species, with only two middle depths (300-700 m) species, the grenadiers, *Macrourus novaezelandiae* and *M. magellanicus* (both >200 000 tonnes in 2003). China’s landings of deepwater fishes are dominated by the largehead hairtail *Trichiurus lepturus* (which accounted for 1.5% of the total world marine fish landings in 2002), and although sometimes listed as a deepwater species, is more correctly a shelf species found in depths <400 m.

Landings of deepwater fishes have risen from <0.5 m tonnes a year in the 1960s to >3 m tonnes by the late 1990s, with more than half of the annual catch taken from the
Seamounts, steep sided undersea mountains, are widely distributed in the world's oceans and usually associated with volcanic activity (Rogers, 1994). Some definitions describe seamounts as features with an elevation greater than 1 000 m, but in practice “seamount” is applied to knolls (elevation 500-1 000 m) and hills (elevation <500 m) that contrast with the surrounding seafloor. Seamounts often occur in clusters along ridges leading to island groups or chains that are physically isolated from other island chains. Seamounts (and oceanic islands) enhance productivity, due to Taylor columns and upwelling of nutrient rich water (Rogers, 1994), and provide a unique deep-sea environment for fishes and invertebrates that are not found in the open ocean (Boehlert and Mundy, 1993; Koslow et al., 2000; Richer-de-Forges et al., 2000). Several teleosts spawn above seamounts where they form dense seasonal aggregations (Koslow et al., 2000).

A high degree of endemism has been reported for benthic invertebrates (~30%) and fishes (12%) on seamounts (Wilson and Kaufman, 1987; Richer-de-Forges et al., 2000; Froese and Sampang, 2004). The macro invertebrates on seamounts tend to be dominated by suspension feeding corals (Rogers, 1994) which are most abundant along the sides and ridges of seamounts and provide habitat for a diverse facultative fauna (Jensen and Frederikesen, 1992). Recent exploration using acoustics and submersibles has revealed unexpectedly widespread and diverse coral ecosystems in deepwaters on continental shelves, slopes, seamounts, and ridge systems around the world (Roberts et al., 2006). In the New Zealand Exclusive Economic Zone macroinvertebrates in trawl samples are made up of Cnidaria (black corals, hard corals, and sea fans), Echinodermata (starfish, sea lilies, and brittlestarfish), Arthropoda (stone crabs and true crabs), and Mollusca (gastropods, octopus and squid); but the greatest invertebrate biomass has been corals (Probert et al., 1997). These large epibenthic organisms are vulnerable to trawling, and corals have been the dominant bycatch in the development of trawl fisheries on some newly discovered seamounts (Anderson and Clark, 2003). On Tasmanian seamounts major impacts were recorded within a few years of the development of the orange roughy Hoplostethus atlanticus fishery; on heavily trawled seamounts (>1 000 trawls) reef aggregate was removed or reduced to rubble, and the invertebrate biomass was 83% lower than on lightly fished seamounts (Koslow and Gwollett-Holmes, 1998). The recovery of these deep-sea corals that may live for centuries (Andrews et al., 2005) is likely to be extremely slow.

Fisheries for teleosts and to a lesser extent for crustacea occur on and around seamounts in the North Pacific Ocean along the southern Emperor and northern Hawaiian Ridge in the North Pacific, in the southwest Pacific Ocean around New Zealand, New Caledonia and Tasmania, and the SE Pacific off Chile, in the North Atlantic around the Azores and on the Mid Atlantic Ridge, and in the South Atlantic. Fisheries have expanded for toothfish around sub Antarctic islands and seamounts in the Southern Ocean and for orange roughy and alfonsino on the Southwest Indian Ocean Ridge.
Atlantic Ocean, but still accounts for only ~5% of the total fish catch. The landing statistics are likely to be under estimates due to illegal, unreported and unregulated (IUU) fishing operations, and discards of bycatch species.

Currently discarded fish waste from processing is used for low-value products such as fish-oils, meals, pet foods, and silage. Bioactive compounds may be extracted from leftover fish-frames, internal organs, and invertebrate bycatch species for biotechnological and pharmaceutical applications, offering the opportunity to add value to fisheries. Some compounds derived from fish waste have been identified as potential nutraceuticals (Kim, S-K. and Mendis, E. 2006). Marine invertebrates that occur around hydrothermal vents may provide enzymes and biochemicals for the biotechnology industries and become target species in the future, raising further issues over exploitation of specialised deepwater habitats.

**Deepwater fisheries and genetic resources**

Aquatic genetic resources have been defined by the 1993 UN Convention on Biological Diversity (CBD) as genetic material of actual or potential value. It has been assumed that such a broad definition encompasses the sum total of all aquatic plants and animals on the planet and that aquatic biodiversity and aquatic genetic resources are almost synonymous terms (Bartley and Pullin, 1999; Pullin, 2000). Fish stocks and bycatch that are exploited, or potentially exploited, by fisheries are all considered as genetic resources (Bartley and Pullin, 1999). However, unlike the terrestrial genetic resources based on plants (PGR), livestock (AnGR), and even aquaculture genetic resources (FiGR, after Pullin, 2000), the deep-sea genetic resources are based on capture from natural ecosystems.

Much of fisheries management has been and continues to be directed towards population and ecosystems management and incorporates genetic resources by default (FAO, 2005b).

The specific application of genetic tools in the management of capture fisheries has been limited to stock identification. Awareness is growing of the genetic structure of fish stocks and the impact of fishing on genetic diversity (Pullin, 2000), but for many marine species the patterns of genetic diversity are poorly understood. The short term pragmatic stock assessment goals to estimate maximum sustainable yields, by necessity have overlooked the long term goals of conserving genetic diversity.

**Deepwater fishes**

Deepwater fishes comprise three major groups: pelagic fish living largely in midwater, with no dependence on the bottom; demersal fish, living close to and depending on the bottom; and benthopelagic fish, living close to the bottom but undertaking vertical migrations in the water mass (e.g. for feeding). Much remains unknown about the biology and distribution of deepwater fishes and new species continue to be discovered (Roberts and Paulin, 1997; Roberts et al., in press). Species exploited by deepwater fisheries include both shelf species, that extend down the continental slopes, and species restricted to depths >400-500 m, and have been grouped into those that aggregate on seamounts and ridges and those more generally dispersed on the continental slope (see Box 2). Many of the commercially targeted species are widespread horizontally, but zoned by depth (e.g. alfonsino and orange roughy), and exhibit specialist adaptations for dispersal and recruitment (Boehlert and Mundy, 1993).

Species diversity is high in the deep-sea and many fishes exhibit unique adaptations, such as bioluminescent organs, modified swim bladders, jaws, and eyes for the deep-sea environment. FishBase lists 1276 bathypelagic species and 2103 bathydemersal species; 798 species of fish have been classified as seamount species (Froese and Sampang, 2004; Morato et al., 2006a). Not all deepwater fishes are well described and molecular tools
Species associated with seamounts

The orange roughy *Hoplostethus atlanticus* has a wide distribution in the Atlantic, Indian and South Pacific Oceans, where it is found between 500-1 800 m, but is most abundant from 750 to 1 100 m. Fisheries have developed around New Zealand, south-east Australia, in the southwest Indian Ocean, off Namibia, Chile, and on the Mid Atlantic Ridge, but the bulk of the catch has been made in the southern hemisphere. The New Zealand fisheries initially concentrated on flat bottom and slope edges, but technical developments, such as GPS navigation, net monitoring, and swathe mapping, coupled with increasing experience in the deepwater fisheries lead to the targeting of orange roughy spawning aggregations on seamounts. The catch of orange roughy from seamounts has increased from about 30% of the total catch in 1985 to 80% by 1995 and has stabilized at 60-70% (Clark, 1999; Clark, and O’Driscoll, 2001).

In spite of the extensive distribution, adults are not highly migratory and movement, inferred from seasonal catches and changes in distribution, is only hundreds of kilometres. Orange roughy eggs remain in the plankton for only about 10 days before descending and hatching near the bottom and the larvae are assumed to be epibenthic (Zeldis et al., 1994). Relatively few juveniles (<1 000) have been caught in bottom trawls around New Zealand (Mace et al., 1990) where the fishery peaked at more than 50 000 tonnes a year (Annala et al., 2000). The species is slow-growing, reaching maturity at 25-30 years of age, and may live for more than 100 years (Smith et al., 1995).

The Oreosomatids, the black oreo (*Allocyttus niger*) and the smooth oreo (*Pseudocyttus maculatus*), support fisheries in the New Zealand and Australian EEZs. In the New Zealand EEZ black and smooth oreo, together with the less abundant spiky oreo (*Neocyttus rhomboidalis*), have been managed under a combined quota. The proportion of oreo catch derived from seamount fisheries increased from ~20% in the 1980s to 65% in the 1990s. Oreos aggregate in the mid slope region and above seamounts at 600-1800 m, and are long lived with estimated maximum ages of 86 years for *P. maculatus* and 150 years for *A. niger* (Doonan et al., 1995). Smooth oreo adults are generally found north of 52° S, but most of the few recorded juveniles have been found between 60 and 68° S (James et al., 1988); only 23 black oreo juveniles have been recorded from the New Zealand EEZ (McMillan, NIWA, unpub.obs.), despite annual catches >25 000 t within the New Zealand EEZ (Annala et al., 2000). Juveniles of both black and smooth oreos are pelagic (James et al., 1988) and settle at approximately 4 and 6 years respectively. The pelagic features and their low Δ14C levels were interpreted as indicating a high latitude origin for black and smooth oreo juveniles (Morison et al., 1999). It is possible that there are single genetic stocks of both species, and that juveniles recruit northwards, after which they show little dispersal and may form discrete ecological stocks.

Alfonsino (*Beryx splendidens*) has a wide distribution in tropical and temperate waters of the Atlantic, Indian and Pacific Oceans and the Mediterranean Sea (Kotylar 1996), and occurs over seamounts and the continental slope in depths between 25–1300 m, but is most abundant between 300–500 m. Maximum age is 20 years and age at maturity is from 6 to 8 years. The adults do not appear to make extensive adult migration to spawning areas (Lehodey et al., 1997), but the larvae and juveniles disperse widely in the pelagic environment for several months before settling on shallow seamounts (Boehler and Sasaki, 1993). *B. decadactylus* has a wide distribution in tropical and temperate waters of except the eastern Pacific Ocean (Kotylar 1996), and occurs on the continental slope and ridges. Relative proportions of *B. splendidens* and *B. decadactylus* are unknown.
Black cardinalfish (*Epigonus telescopus*) is widely distributed in the North Atlantic from Iceland to the Canary Islands, in the western Mediterranean, and in the South Atlantic, Indian, and southwest Pacific Oceans. The species occurs between 200–1400 m but is most common between 600–900 m. The juveniles are pelagic and undergo major ontogenetic changes; little is known of adult movements. Unvalidated otolith readings indicate slow growth and longevity, with maximum ages ~100 years. Around half the New Zealand catch (~2 000 tonnes year) has been taken as bycatch, with 80% taken in the orange roughy fisheries. In the North Atlantic black cardinalfish are taken as bycatch in trawl and long-line fisheries (Pineiro et al., 2001).

Toothfish are large notothenoids living in Antarctic and sub-Antarctic waters. The two species are circumpolar, the *Antarctic toothfish* (*Dissostichus mawsoni*) is found at high latitudes south of the Antarctic Convergence around 60° S, while the *Patagonian toothfish* (*D. eleginoides*) ranges from about 50° S to 65° S, around sub Antarctic Islands and seamounts, between 50-60° S, and on the Patagonian Shelf and the southern coast of Chile to 30°S (Gon and Heemstra, 1990). *D. eleginoides* reaches a large size >200 cm and age of 50 years (Horn, P. 2002). It is targeted by trawl and long-line fisheries between 70 - 1 800m, and was lightly exploited until the mid 1980s, with catches around several hundred tonnes a year. Catches increased rapidly during the 1990s; unofficial estimates suggested that catches reached more than 80 000 tonnes in 1996-97, with large IUU fishing activities (ISOFISH 1998), which have subsequently declined following the introduction of a catch documentation scheme. *D. mawsoni* reach a length of 175 cm and age of ~35 years (Horn, 2002) and have become the target of a number of new and exploratory fisheries since the mid 1990s, with TACs determined by CCAMLR.

Large catches of the *pelagic armourhead* (*Pseudopentaceros wheeleri*) were taken from seamounts (with summits 250-600 m) along the Emperor-Northern Hawaii Ridge in the central North Pacific during the 1970s. Annual catches were estimated at 50 000-200 000 tonnes, but were reduced to a few thousand tonnes in the late 1970s (Boehlert, 1986; Somerton and Kikkawa, 1992). The species is fast growing with a long pelagic juvenile phase and maximum age of 4 years.

*Roundnose grenadier* (*Coryphaenoides rupestris*) is abundant in the North Atlantic north of 50° N, from Newfoundland Banks to Rockall at 600-800 m, and occurs down to 2 000 m; long lived > 60 years; matures at agea 8-10 years. *C. rupestris* is caught in mixed trawl fisheries with black scabbard fish in the NE Atlantic. Geographically distinct populations exist on the Mid-Atlantic ridge and the Hatton Bank, but its genetic relationships are unknown. A fishery developed in the Northeast Atlantic in the mid 1970s peaked at ~80 000 tonnes and declined rapidly to ~6 000 tonne a year by 1980; the fishery began in the north of the range and moved southwards in the NW Atlantic (Atkinson, 1995). *C. rupestris* has recently been identified as critically endangered (Devine et al., 2006).

*Sebastes* spp. (*redfish* and *ocean perch*) have supported the longest deepwater fisheries in both the Atlantic and Pacific Oceans. Many species are long lived (up to 100 years), with slow growth rates and late maturity (> 20 years). *Sebastes alutus* is found <50–825 m in the North Pacific, from Honshu, Japan through the Bering Sea to California, with a maximum age of 100 years (Leaman, 1991). The primary focus of deepwater fisheries has been on the upper slope off North America, with catches peaking in the 1960s at around 450 000 tonnes, and since fluctuating at 5 000-30 000 tonnes (Ianelli and Zimmerman, 1998). The fishery has extended into deepwater and exploits several other species of scorpaenids (Ianelli and Zimmerman, 1998)

The *Sebastes* fishery in the Northwest Atlantic is based on the redfish complex *S. fasciatus*, *S. mentella*, and *S. marinus*, which are caught on the shelf edge and the upper
Continental slope species

The ling (*Molva molva*) is found in the Northwest and Northeast Atlantic on the continental shelf, and is common between 100-400 m; it is found down to 1000 m. The only significant fisheries are in the Northeast Atlantic, where landings have been around 50-60 000 tonnes since the 1970s, but recent CPUE data indicate severe depletion (ICES, 2005a). Ling reach 30 years of age and grow to 200 cm in length.

The blue ling (*Molva dypterygia*) is common between 350-500 m, and ranges between 150-1 000 m in the Northeast and Northwest Atlantic. Landings peaked at 35 000 tonnes in the 1980s, but declined to <10 000 tonnes. It reaches 20 years of age and 155 cm in length. Recent CPUE data indicate a severe depletion. Its growth rate is unknown (ICES, 2005a).

The tusk (*Bromse bromse*) is found in the Northwest and Northeast Atlantic on the continental shelf to 1 000 m, but the only fisheries are in the Northeast Atlantic. It reaches a maximum size 120 cm and a reported age of 20 years. Its landings are in decline and CPUE indicates a severe depletion. Its growth rates are unknown (ICES, 2005a).

Hoki or blue grenadier (*Macruronus novaezelandiae*) support the largest fishery in the New Zealand EEZ and are caught between 300-700 m. It occurs from 10-900 m, matures at 4-5 years and has a maximum age of 20-25 years. It is found around New Zealand and Tasmania. Annual catches in New Zealand peaked at 269 000 tonnes in 1997-98; the current TAC has been reduced to 100 000 tonnes. The whiptail hake (*Macruronus magellanicus*) supports trawl fisheries in the South Atlantic and South Pacific around South America from Punta Medanos Argentina to Valparaíso Chile. It is caught by the purse-seine fleet off central-south Chile. It reaches maximum age of ~20 years.

Greenland halibut (*Reinhardtius hippoglossoides*) is found in the North Pacific and North Atlantic Ocean on the shelf down to 2000 m; it reaches a maximum age of 30 years and length of 120 cm. The Northwest Atlantic fishery for *R. hippoglossoides* remained high over the 1960s to 1990s, but grew rapidly during the early 1990s with the entry of Spanish vessels into the fishery, leading to the much publicised Canada-Spain “turbot war” in 1993. The mean size of fish has declined rapidly and the bulk of the catch is made up of fish smaller than the size at maturity. A major collapse of the fishery appears likely (Haedrich *et al.*, 2001).

Black scabbardfish (*Aphanopus carbo*) has a wide distribution in the Northeast Atlantic in 200-1 600 m. There are longline fisheries off Madeira and Portugal and more recently *A. carbo* has become an important species in the mixed bottom-trawl fishery that developed in the Rockall Trough in the 1990s where it is caught with *C. rupestris*. Age estimates vary from 8-25 years with a maximum length of 110 cm. The stock composition of this species is unknown, but element composition of its otoliths indicates differences between the northern and southern areas of the Mid-Atlantic Ridge (Swan *et al.*, 2003). The eggs, larvae and small juveniles are unknown.

The sablefish (*Anoplopoma fimbria*) is found in the North Pacific: Bering Sea coasts of Kamchatka, Russia and Alaska southward to southern Japan and central Baja California.
It supports major fisheries in the NE and NW Pacific and reaches a maximum size of 200 cm, and maximum reported age of 114 years. It is found from the surface to 2 700 m: the juveniles are pelagic and migratory.

Deepwater sharks
In the Northeast Atlantic 12 species of shark are caught in the deepwater fisheries. The wide distribution of deepwater sharks means that fishers in deepwater areas cannot avoid catching them and the catch has risen from <100 t a year in the late 1980s to ~11 000 tonnes by 2003. The Portuguese dogfish (Centroscymnus coelolepis) and the leafscale gulper shark (Centrophorus squamosus) are target species in long-line and gill net fisheries but they, and other sharks, are taken as bycatch in trawl fisheries targeting orange roughy, roundnose grenadier, blue ling and longline fisheries for black scabbard fish. As trawlers have started to fish further down the continental slope the species mix of sharks has changed from that dominated by leafscale gulper shark in the early stages of the fishery to the Portuguese dogfish (ICES 2005b); more than 95% of the ICES shark catch probably consists of these two species; the other shark species have low commercial value and are discarded. A combined TAC applies to all deepwater shark species for 2006-07 (ICES, 2005b). The lack of species-specific catch data may have disguised an extreme decline of vulnerable species. Although it has been difficult to advise on a sustainable catch with limited information, the current fishing effort in the North Atlantic is recognised as too high (ICES, 2005b).

Both C. squamosus and C. coelolepis have wide distributions and depth ranges. C. squamosus is found between 150–2 400 m in the Eastern Atlantic, the Western Indian Ocean and Western Pacific, and although caught on the bottom 600–1 000 m it is pelagic above deepwater (4 000 m). It reaches a maximum age of 60–70 years and size of 160 cm. C. coelolepis is caught between 500-1 500 m and are found between 270-3 600 m in the Western and Eastern Atlantic, the Mediterranean and Western Pacific. It reaches a maximum size of 120 cm; Its maximum age is unknown, and it is near threatened.¹

Deepwater sharks in other fisheries are vulnerable, whether taken as targeted species or as bycatch. The dumb gulper shark (Centrophorus harrissoni), which is possibly restricted to Western Australia, Tasmania and New South Wales and caught in a deepwater trawl fishery off New South Wales, is critically endangered (FishBase).

¹ A general term used to cover taxa whose survival is uncertain (FishBase).

are being used to resolve taxonomic questions of species identity. DNA barcoding initiatives (Hebert et al., 2003) will provide tools for the rapid identification of species in processed products.

Some of the commercially important deepwater fishes exhibit extreme life history traits with slow growth rates, high longevity (~100 years) and late maturity (15-25 years), see Box 2. In addition some species appear to exhibit long periods of low recruitment (Koslow et al., 2000). For species with episodic recruitment the removal of older fishes may reduce the ability of populations to withstand extended periods of very low recruitment (Koslow et al., 2000); for example the Pacific Ocean perch (Sebastes alutus) in lightly and heavily fished populations shows 73% and 7% respectively of fish older than 20 years (Leaman, B.M. 1991).
4. STATUS AND TRENDS OF THE GENETIC RESOURCES

Genetic resources and stock structure
A knowledge of the stock structure of marine fishes is important for the management and conservation of genetic resources. Several approaches are used to measure relationships among spatially isolated populations, most are based on ecological measures, such as a parasite load (McKenzie, 2002) or accumulation of trace elements (Thresher, 1999), or environmentally sensitive characters such as morphometrics and meristics (Cadrin, 2000). Molecular tools provide an alternative, indirect measure of dispersal and gene flow. Genetic diversity measured with most molecular methods is assumed to be selectively neutral and non-adaptive with respect to fitness. In general marine fishes have higher levels of genetic diversity than anadromous species, which in turn have higher levels than freshwater species (Gyllenstein, 1985; Ward et al., 1994), a trend that probably results from larger evolutionary effective population sizes in marine fishes (Dewoody and Avise, 2000).

Marine fishes on average show less spatial genetic differentiation than anadromous and freshwater species, due to the fewer barriers to gene flow in the marine environment. Marine dispersal is constrained by the length of time of the pelagic larval and juvenile stages, by behavioural mechanisms, and by physical barriers such as gyres and ocean fronts. An inverse relationship has been reported between genetic differentiation and dispersal potential in small shelf fishes (Waples, 1987; Doherty et al., 1995), but many of the larger shelf fishes show little genetic differentiation over ocean wide scales (Hauser, and Ward, 1998), possibly due to a combination of large population sizes and high mobility.

Deepwater species with potential for extensive dispersal through pelagic juvenile stages, such as the Pacific armourhead, *Pseudopentaceros wheeleri* (Martin et al., 1992), the alfonsino *Beryx splendens* (Hoarau and Borsa, 2000; Aboim, 2005), the wreckfish *Polyprion americanus* (Sedberry et al., 1996; Ball et al., 2000), and the silver roughy *Hoplostethus mediterraneus* (Smith unpublished observations) exhibit ocean-wide genetic population structures. In the wreckfish, microsatellite allele frequencies were homogeneous in the eastern and western North Atlantic and Mediterranean, but heterogeneous between the North and South Atlantic Ocean (Ball et al., 2000), a genetic discontinuity supported by differences in mitochondrial (mt) DNA haplotype frequencies; and by implication there is little contemporary gene flow across the tropics (Sedberry et al., 1996). In the redfish *Sebastes mentella* genetic homogeneity observed over 6 000 km probably results from larval gene flow in the cyclonic circulation of the central North Atlantic, although at wider spatial scales there is evidence for three genetically differentiated groups around the Gulf of St Lawrence/Newfoundland; the Grand Banks to the Faroes; and the eastern Atlantic and the Barents Sea (Roques et al., 2002). In the hoki *Macruronus novaizelandiae* there is no genetic differentiation among spawning stocks within the New Zealand EEZ (Smith et al., 1996), but there is genetic differentiation across the Tasman Sea (Milton and Shaklee, 1987).

Likewise for the black oreo *Allocyttus niger* and smooth oreo *Pseudocyttus maculatus* which have extensive pelagic dispersal during the juvenile stages, no significant genetic differentiation was found among black oreo and among smooth oreo samples from the New Zealand EEZ (Smith et al., 2002). At wider scales a lack of genetic differentiation was reported in smooth oreo samples from Western Australia, Tasmania, and New Zealand, with a different suite of allozyme and mtDNA markers, and little evidence for genetic differentiation between black oreo samples from New Zealand and Tasmania (Ward et al., 1998).

Mitochondrial DNA haplotype data indicated a strong genetic differentiation between populations of the viviparous blackbelly rosefish *Helicolenus dactylopterus* from the NE and NW Atlantic Ocean (Aboim, 2005; Aboim et al., 2005). The
application of microsatellite DNA markers has revealed finer population structure within the central NE Atlantic Ocean off Portugal and around the Azores archipelago (Aboim, 2005).

Contrasting patterns of genetic differentiation have been reported in some congeneric pairs of species. In the Patagonian toothfish (D. eleginoides) a study of allozyme markers showed no significant regional differentiation among samples from the Southern Ocean, while microsatellite DNA markers showed significant heterogeneity, rejecting the null hypothesis of a single stock (Smith and McVeagh, 2000). Samples from the Indian Ocean were homogeneous for both mitochondrial DNA and microsatellite markers (Appleyard and Williams, 2004), but heterogeneity was found among samples from Macquarie Island, Heard and MacDonald Islands (Appleyard et al., 2002). A major genetic break has been reported north and south of the convergence zone in the Atlantic Ocean (Smith and Gaffney, 2000a; 2000b; Shaw et al., 2004). Recent studies with single nucleotide polymorphisms (SNPs) in D. eleginoides have revealed finer population structure with differentiation within ocean basins (Gaffney, University of Delaware pers.com.). In contrast, populations of the Antarctic toothfish (D. mawsoni) appear to be characterized by very low mitochondrial sequence diversity, and homogeneous frequencies of nuclear alleles and mitochondrial haplotypes among sea areas. These preliminary genetic data provide little support for the hypothesis of separate regional stocks of (D. mawsoni) (Smith and Gaffney, 2005).

In the congeneric alfonsinos (Beryx splendens) and (B. decadactylus) analyses of mtDNA haplotype data revealed major differences in the structure and history of the populations of the two species. B. splendens appears to have one population in the Northeast Atlantic, while B. decadactylus, exhibits lower genetic diversity but strong genetic differentiation between Cape Verde and the other populations in the NE Atlantic Ocean (Aboim, 2005).

The orange roughy (Hoplostethus atlanticus) and silver roughy (H. mediterraneus) have wide distributions in the North Atlantic, Indian and Pacific Oceans, but different dispersal potentials. Orange roughy have weak dispersal potential: their pelagic eggs sink and hatch near the bottom (Zeldis et al., 1994) and exhibit genetic differentiation at small spatial scales in the SW Pacific Ocean and Tasman Sea (Smolenski et al., 1993; Smith et al., 1996; Smith and Benson, 1997; Smith et al., 1997). In contrast silver roughy have a long pelagic phase and show little genetic differentiation at the oceanic scale (Smith, unpublished results). Local declines among orange roughy fisheries on neighbouring seamounts suggest that they may be independent units in the ecological time frame of fisheries management, in the absence of detectable genetic differentiation.

Genetic diversity in deep-sea soft sediment invertebrates and deep-sea corals

The deep-sea soft-sediment environment hosts a diverse and often highly endemic fauna of uncertain origins. Little is known of the genetic resources and the impact of trawling on these soft substrates, but some broadly distributed invertebrates exhibit genetically divergent populations in the absence of morphological divergence, and may represent cryptic species (Etter, 1999; Zardus et al., 2006). High levels of genetic diversity were found in the protobranch bivalve (Deminticula atacellana), which is widespread throughout the Atlantic Ocean in soft sediments at bathyal and abyssal depths. Samples from localities in the North American, West European and Argentine basins were divided into four major clades, with DNA haplotypes unique to each basin (Zardus et al., 2006). Genetic divergence was greater among populations at different depths within basins, than among those at similar depths in separate basins, indicating population differentiation at small (100s kms) spatial scales (Chase et al., 1998; Zardus et al., 2006). Depth-related divergence has also been reported in the deepwater amphipod (Eurythenes gryllus) (Bucklin et al., 1987; France and Kocher, 1996), and this general finding may reflect historical patterns of colonization, or strong environmental
selective gradients, or horizontal dispersal in the deep-sea (Bucklin et al., 1987; Etter et al., 2005; Zardus et al., 2006).

There have been few genetic studies on deepwater corals to identify dispersal and connectivity among seamounts (Baco et al., 2006); yet such data are required for the development of management strategies for these species that dominate the fragile ecosystems. Low-sequence divergences were found among some deep-sea octocorals, but preliminary results for the bamboo corals (Keratoisidinae) in the SW Pacific suggest that some species are widespread and are not restricted to seamounts (France, and Hoover, 2002; Smith et al., 2003). Genetic data for the scleractinian coral (*Lophelia pertusa*), the main framework-building species in the Northeast Atlantic at depths between 200 and 1 000 m, have revealed distinct offshore and fjord populations. The levels of genetic diversity in *L. pertusa*, and the contribution of asexual reproduction to the maintenance of the subpopulations were highly variable among sites (Goff-Vitry et al., 2004).

Potential loss of genetic diversity in small populations

The genetic composition of a population can change over generations due to random events. Changes due to genetic drift are most likely in small populations and are expected to be weak in marine fishes with large populations (*N* >10^7). However, sweepstake events driven by very high larval mortalities can reduce *N*, (the number of individuals contributing to the next generation) by several orders of magnitude from the census population size, *N* (Hedgecock, 1994). Estimates of *N*, in several shelf fishes are considerably smaller then census sizes (Bagley et al., 1999; Chapman et al., 2002; Hauser et al., 2002; Turner et al., 2002). Long-lived fishes with overlapping generations and annual spawning events are protected from loss of genetic variability due to drift, through the “storage effect” of year classes that buffer annual decreases in *N*, (Gaggiotti and Vetter, 1999). However, additional pressures from fishing practices that lead to population declines and loss of juvenile habitat, imposed on sweepstake recruitment events, may lead to loss of genetic diversity (Chapman et al., 1999b; Hauser et al., 2002). Temporal genetic variation maybe enhanced in deepwater fishes, because the low productivity environment may restrict individual fish from spawning annually, leading to low and patchy recruitment (Leaman and Beamish, 1984).

Within-area temporal variation has been reported in orange roughy (Smolenski et al., 1993; Smith and Benson, 1997); grouper (*Mycteroperca microlepis*) (Chapman et al., 1999b); and hake (*Merluccius merluccius*) (Lundy et al., 2000); and may result from stochastic events in progeny survival (Chapman et al., 1999a). The low *N,/N* ratios demonstrated in several shelf species are equally likely to occur in some deepwater species, and potentially lead to loss of genetic diversity in collapsed stocks.

Genetic diversity and selective fisheries

Substantial changes in life history traits, in particular age and size at maturity, have been reported in heavily exploited stocks on the continental shelves in the North Atlantic (Smith, 1994; Stokes and Law, 2000; Dieckmann et al., 2006 in press). These changes may result from environmental change, the direct selective effects of fishing, or a compensatory response to reduced stock densities (Law, 2000). The compensatory response to a reduction in stock size promotes growth rate, resulting in a decrease in the age at maturity but an increase in the size at maturity, and may conceal long-term selection effects that would favour early maturing genotypes (Roche, 1998). These responses may be non-exclusive making it difficult to untangle the compensatory and evolutionary components of these observed changes (Law, 2000).

Reaction norms for age and size at maturation have been used to estimate the probability of maturing at each relevant age and size, and thereby separate the genetically determined character from the plasticity in maturation that results
from changes in growth rate (Heino et al., 2002; Engelhard and Heino, 2004). Growth-related phenotypic plasticity appears to have been largely responsible for recorded changes in early maturity in the Norwegian spring herring \textit{(Clupea harengus)} (Engelhard and Heino, 2004), but in Atlantic cod \textit{(Gadus morhua)} (Heino, 2002) and plaice \textit{(Pleuronectes platessa)} (Rijnsdorp et al., 2005) evolutionary changes appear to have occurred in response to heavy fishing (Law, 2000). These evolutionary changes are supported by controlled selection experiments that have demonstrated that fisheries have the potential to cause rapid evolution in life history traits (Conover, 1998; Conover et al., 2005). Life history traits in populations of Atlantic cod off southern Labrador and eastern Newfoundland continually shifted towards maturation at earlier ages and smaller sizes, before the fishery finally collapsed in the 1990s. These changes in life history could provide a tool to give warning signals before more overt changes occur to populations (Olsen et al., 2004).

Directional selection, through size-selective harvesting has not been demonstrated in deepwater fishes, in part due to the lack of long term data sets and, for long-lived species the long response time of the population to fishing. A number of changes were observed in a major orange roughy fishery on the Chatham Rise (New Zealand) over a 19-year period. The species distribution showed a marked contraction, with aggregations becoming centred around seamounts, or localised areas of the slope, and the biomass declined substantially to about 20% of virgin levels (Clark et al., 2000). However size structure and size or age at maturity did not change markedly over the same period. Biological changes may not have been apparent because orange roughy is a long-lived, slow-growing species, with low productivity (Clark et al., 2000).

Deepwater fisheries have only been operating in the Northwest Atlantic Ocean since the 1970s, but already four out of five species (the roundnose grenadier \textit{(Coryphaenoides rupestris)}, the onion-eye grenadier \textit{(Macrourus berglax)}, the blue hake \textit{(Antimora rostrata)}, and the spinytail skate \textit{(Bathyraja spinicauda)}) have declined by 25-57% in mean size over 17 year period, so that fewer fish reach maturity and breed (Devine et al., 2006). In the Northwest Atlantic fishery for Greenland halibut \textit{(R. hippoglossoides)} the mean size of fish has declined rapidly and the bulk of the catch is made up of fish smaller than the size at maturity, and a major collapse of the fishery is likely (Haedrich et al., 2001).

**Endangered species**

Only a few marine fishes have been listed as endangered (although the list is growing) and fewer appear to be close to extinction, e.g. skates (Brander, 1981; Casey and Myers, 1998). Traits of several deepwater species, such as long-life span, large body size, low natural mortality and late sexual maturity, are likely to make them more vulnerable to extinction than shelf species, in particular species that aggregate above seamounts. A review of the extinction risk in marine fish found that large body size and late maturity were the best predictors of vulnerability to fishing; there was no evidence that high fecundity conferred increased resilience (Reynolds et al., 2005). Much of the evidence for extinction risk comes from shelf species, where inshore sub-populations of Atlantic cod and herring have been driven to extinction or have had insufficient time to recover from severe depletions (Smedbol and Stephenson, 2001).

Different criteria have been used to express the risk of extinction. The widely used IUCN system uses the graded terms vulnerable, endangered, and critically endangered and is applied to all organisms regardless of life history strategy. The IUCN criteria may overestimate the extinction risk for many marine fishes with their high intrinsic rates of increase, and for which management plans allow for stock biomass targets of 20-30% of the virgin biomass (Musick, 1999). Stocks of some pelagic shelf fishes have collapsed with severe reductions (1/3000) in population size, but have shown evidence of recovery. In general these species are characterized by small body size and early
maturity, the converse of traits that were the best predictors of vulnerability to fishing (Reynolds et al., 2005). Pelagic species showing the greatest declines have shown the slowest recoveries (Beverton, 1990). The American Fisheries Society (AFS) developed a precautionary set of criteria to predict the risk of extinction in marine fishes, which includes rarity, specialization in habitat requirements, endemism or small range, and population decline, and also aims to recognise distinct population segments (DPS) when data are available (Musick, 1999). Rare species, because of evolutionary or ecological factors or crypsis, would be classified as vulnerable until further data were available. Species that are endemic or occur over a small range where the habitat is under threat from degradation would be classified as vulnerable, and where habitat loss has occurred they would be classified as endangered or threatened. Species with specialised habitat requirements, but that occur over wide geographic ranges, may also be vulnerable when the specialised habitat is subject to degradation or destruction (Musick, 1999). However the lack of knowledge about critical minimum population size and possibility of depensation create the greatest problems in assessing the extinction risk in marine fishes (Musick, 1999). Consequently the AFS proposed evaluating the resilience of the DPS using four levels of productivity: high, medium, low, and very low. Fish with late maturity (5–10 years), a long life span (>30 years), and high fecundity (>10⁴), typical of some deepwater fishes (and some show even later maturity and greater longevity), would be classified as very low productivity and would have a lower threshold to extinction than a species with medium or high productivity (Musick, 1999). Non-target species, that include teleosts endemic to seamount complexes and elasmobranchs with very low productivity, are likely to be vulnerable. For the few seamount fishes for which there are adequate biological data, most species have a low or very low productivity, and low resilience to exploitation (Froese and Sampang, 2004). It has been predicted that more seamount populations will be depleted and some will be extirparted if fishing continues at current levels (Morato et al., 2006a).

Five deepwater species (the roundnose grenadier C. rupestris, the onion-eye grenadier M. berglax, the blue hake A. rostrata, the spinytail skate B. spinicauda, and the spiny eel Notocanthus chemnitzi) in Northwest Atlantic fisheries appear to meet the IUCN and AFS criteria of being critically endangered (80% decline in 10 years or three generations, or whichever is longer), showing overall declines in relative abundance of 87–98% in <20 years of exploitation, and higher estimated declines over three generations (Devine et al., 2006). If the IUCN criteria are applied, the Dumb gulper shark (Centrophorus harrissoni) caught in the deepwater fishery of New South Wales is critically endangered; the deepwater bluntnose sixgill shark (Hexanchus griseus), circumglobal in tropical and temperate seas, is vulnerable (20% decline in 10 years or three generations or whichever is longer) while the kitefin shark (Dalatias licha) and the leafscale gulper shark (Centrophorus squamosus) are at lower risk, near threatened (i.e., survival is uncertain), along with several other species of shark in shelf waters (IUCN Red List).

The blue skate (Dipturus batis) is endangered and extirpated by trawling over much of its range in the eastern North Atlantic. The shallow water bocaccio (Sebastes paucispinis) found on seamounts in the eastern Pacific is critically endangered, while the deepwater shortspine thornyhead (Sebastolobus alascanusis) in the North Pacific is endangered. The Atlantic halibut (Hippoglossus hippoglossus) caught in shelf and slope fisheries in the North Atlantic is also endangered (see FishBase).

5. MAJOR INTERNATIONAL INITIATIVES, AGREEMENTS AND INSTRUMENTS

Genetic resources at the species and ecosystem levels are equivalent to ecological resources for which the management issues are well documented in the fisheries literature. The rapid development, and in some cases rapid depletion, of deepwater fishery is of major concern to fisheries managers around the world, and has been
identified repeatedly at the local, regional and international levels (Koslow et al., 2000; Haedrich et al., 2001; Molenaar, 2004; FAO, 2005b; Morato et al., 2006b). ICES have recognised that most exploited deepwater fishes are harvested unsustainably and radical reductions in fleets, in particular bottom trawlers are required to reduce effort and to conserve vulnerable habitats (ICES, 2005a).

Many deepwater fisheries occur in high seas areas adding to the problem of management and regulation. Urgent action is required at the global level, to avoid shifting the deepwater fishing problem from one region to another. Concerns about the apparent inertia in developing and implementing fisheries legislation, especially in the international arena, have lead to NGOs calling for the designation of large-scale protected areas, and for a moratorium on bottom trawling until area management regimes can be implemented. Several countries have small-scale closures for deepwater fisheries within their territorial waters.

1995 UN Fish Stocks Agreement (FSA)
The 1995 UN Fish Stocks Agreement sets out the principles for the conservation and management of straddling fish stocks and highly migratory fish stocks and establishes that management be based on the precautionary approach and the best available scientific information (UN, 1995). The Agreement builds on the fundamental principle, established in the 1982 UN Convention of the Law of the Sea that States should cooperate to ensure conservation and promote the objective of the optimum utilization of fisheries resources both within and beyond the EEZs. The 1995 UN Fish Stocks Agreement was signed by 59 States and entities, but some major fishing nations, such as China and the Republic of Korea have not yet ratified the Agreement. The Agreement does not cover deepwater stocks found exclusively outside the 200 mile EEZs (i.e. discrete high seas stocks), but in practice States have been applying it to discrete high seas stocks (e.g. the South East Atlantic Fisheries Organisation).

A review meeting of the FSA in May 2006 identified a series of actions for States individually, and collectively through regional fisheries management organizations, to ensure the conservation and sustainable use of straddling fish stocks and highly migratory fish stocks, and that these principles should be applied to fish stocks in the high seas.1

1995 FAO Code of Conduct for Responsible Fisheries
The 1995 FAO Code of Conduct for Responsible Fisheries (FAO, 1995) is a voluntary comprehensive instrument that sets out the principles and standards for the conservation and management of all fisheries and aquaculture including processing and trade in fish and fishery products, research and the integration of fisheries and aquaculture into coastal management areas. The Code refers to the role of Regional Fisheries Bodies to establish responsible international fisheries regimes.

Regional fisheries management organizations or arrangements (RFMOs)
There are 44 regional fisheries bodies that cover three categories: RFMOs, Advisory bodies, and Scientific bodies (FAO, Fisheries). Of these, 17 RFMOs are responsible for establishing management measures and some have regulatory powers in their jurisdictions, although many have a purely advisory role. Major problems for the current RFMOs relate to decision making, the allocation of resources to new entrants (principally developing countries that do not have a historical catch record) and the impacts of IUU fishing.2

2 http://www.fao.org/docrep/008/s0098e/s0098e06.htm
Most RFMOs have common responsibilities (Devaney, PL 2005) to:

- collect and distribute fishery statistics,
- provide evaluations of the state of fish stocks in their area of jurisdiction,
- determine the total allowable catch (TAC) quotas,
- set limits on the number of vessels allowed to exploit the fishery,
- control fishing opportunities by RFMO participants using such measures as area and seasonal closures and bycatch limits,
- regulate the types of gear used and conduct inspections to ensure compliance,
- monitor and enforce adherence to the rules of the RFMO and
- oversee the scientific research conducted within the fishery.

Australia, Chile, and New Zealand are promoting the development of a South Pacific Regional Fisheries Management Organisation (SPRFMO) to address governance of high-seas fisheries from the eastern Southern Indian Ocean, across the Tasman Sea and South Pacific Ocean to the Pacific EEZ’s of South America. Several other countries have fishery interests in the area, notably Russian Federation, Ukraine, China, the Republic of Korea and the European Union. Currently there is little or no control over fishing methods or the management of fish stocks, other than for highly migratory tunas, in this extensive region. Other RFMOs, such the Western and Central Pacific Fisheries Convention (WCPFC) and the Inter-American Tropical Tuna Convention (IATTC) cover parts of this region, but their mandates relate only to highly migratory species.

The High Seas Task Force

The High Seas Task Force was established in 2003 to develop an action plan to combat IUU fishing (see below) on the high seas; membership consists of a group of fisheries ministers from Australia, Canada, Chile, Namibia, New Zealand, and the UK; and international NGOs – WWF, the World Conservation Union (IUCN) and the Earth Institute. Although established in 2003 the first report was not released until 2006. The High Seas Task Force aims, inter alia, to develop a Global Information System (GIS) on high seas fishing vessels that will make available information on the characteristics, ownership, and operations of all high seas fishing vessels. The GIS will also identify vessels previously black-listed by RFMOs, with the intention to make it difficult for IUU operators.

The Marine Stewardship Council (MSC) and ecolabelling of fish products

The MSC is an independent non-profit organisation that aims to use consumer purchasing power to enhance responsible management of seafood resources to ensure the sustainability of global fish stocks and the health of the marine ecosystem. The MSC has developed an environmental standard for sustainable and well-managed fisheries and uses a product label to reward responsible fishery management and practices. Certification is only granted if there is consensus amongst independent assessors that the fishery meets the MSC standard. Consumers, concerned about overfishing and its environmental impacts are able to choose seafood products which have been given MSC certification. The New Zealand hoki (Macruronus novaezelandiae) was the first large whitefish and only deepwater fishery to achieve MSC certification, for a period of five years.

Other NGOs have developed eco-labels that rank fish species by the sustainability of the fisheries so as to provide information to consumers. Eco-labels may be in conflict with the MSC, for example the New Zealand Forest and Bird Society’s Best Fish Guide advises consumers to avoid eating hoki, and lists this as a worst choice species, due to the bycatch of fur seals, albatrosses and petrels and management practices in the fishery.

Guidelines for ecolabelling fish products have been developed by the FAO Committee of Fisheries (COFI) for governments and organizations that maintain, or
are establishing, labelling schemes for fish and fishery products from well-managed marine capture fisheries (FAO, 2005a). The general principles for ecolabelling schemes include the need for reliable and independent auditing, transparency of standard-setting and accountability and the need for the standards to be based on good science.

6. KEY SCIENTIFIC AND MANAGEMENT ISSUES TO BE ADDRESSED

Many deepwater species differ from shelf species in that they exhibit high longevity and late maturity; some have extensive but localised distributions in a low productivity environment. However there are limited scientific data available for many species. Working Groups at the Deep-sea 2003 conference identified several scientific areas where additional data are desirable for the management of deepwater species (FAO 2005b):

- accurate catch data,
- time series of abundance,
- stock identity and distribution information,
- life-history information,
- population biology statistics and age-frequency data,
- ability to make use of the most recent developments in fisheries resource management

Fishery subsidies

One component of reducing fishing capacity is the reduction or removal of subsidies (Pauly et al., 2002). In general, the provision of subsidies increases the net returns from fishing and leads to an increased pressure on deepwater fish stocks, although simply removing subsidies will be ineffective in the absence of other management regimes (Cox, 2005). Subsidies, and other incentives such as accelerated depreciation for vessels, that encourage the expansion of capacity, such as vessel construction, may lead to increased pressure on deepwater fisheries that are technology driven. The converse, subsidies for vessel decommissioning, will only be effective if the vessels are scrapped (to avoid transfer to another fishery) and not replaced by new vessels (Cox, 2005). Rising fuel prices might also contribute to the restriction of some deepwater fishing operations.

Illegal, unreported and unregulated (IUU) fishing

The global excess of fishing capacity has contributed to illegal, unreported and unregulated (IUU) fishing and is recognized as a major threat to the long term sustainability of the world’s fish stocks (FAO, 2004a). IUU fishing on the high seas is generally described as any fishing that takes place within the jurisdiction of a RFMO, but not in compliance with its regulations. IUU fishing is more broadly defined by the FAO as fishing activities in the area of application of a relevant RFMO that are conducted by vessels without nationality, vessels flagged to a State not party to that organization, or by a fishing entity, in a manner that is not consistent with, or contravenes, the conservation and management measures of that organization (FAO 2004a).

A number of measures aimed at combating IUU fishing, have been adopted by States and RFMOs, but despite these measures IUU fishing appears to continue because mobile fishing fleets are able to move rapidly between areas. FAO have developed a voluntary instrument within the framework of the Code of Conduct, the 2001 FAO International Plan of Action to Prevent, Deter and Eliminate Illegal, Unreported, and Unregulated Fishing (IPOA-IUU).³

³ http://www.fao.org/docrep/005/y3554e/y3554e00.HTM
IUU fishing for toothfish has consisted largely of illegal fishing within the EEZs of sub-Antarctic island territories within the CCAMLR Area and unregulated and unreported fishing both within and outside the CCAMLR Area. Most IUU fishing is thought to have occurred in the Indian Ocean sector around Crozet, Heard, Kerguelen and Prince Edward Islands. Increased surveillance activity in these areas has forced illegal operators to more remote areas, such as the waters around Ob and Lena Banks, and the waters around South Georgia.4

CCAMLR introduced a catch documentation scheme (CDS)5 that became binding on its members in May 2000. The Scheme tracks the landings and trade flows of toothfish caught in the CCAMLR Area by requiring landings and trans-shipments of toothfish to be accompanied by a valid CCAMLR Catch Document. The CDS identifies the origin of toothfish entering the markets of all participants in the Scheme. CDSs are a promising tool for other RFMOs to encourage legal fishing, and certify that fish entering the markets were caught in compliance with regional fishing regulations. However IUU fishers who land their catch into non-member port states will continue to undermine conservation measures.

In addition, CCAMLR maintains a list of vessels with a history of IUU fishing and has passed a resolution to avoid flagging and licensing non-Contracting Party vessels to fish in CCAMLR waters when the vessels have a history of IUU fishing. Several RFMOs have also instituted a system of black listing IUU fishers as a cost effective enforcement tool and prohibit black listed vessels from landing their catches in member ports.

New technologies are creating opportunities for RFMOs to better monitor vessels and catches and applications are supported by the High Seas Task Force. Vessel Monitoring Systems (VMSs) installed on fishing vessels allow RFMOs to receive up-to-the-minute data on the locations of member vessels. DNA barcoding of marine fish will provide tools for the rapid identification of species in processed products.

Protected areas
The international community, including NGOs, have expressed concern over the loss of macro invertebrates taken as bycatch in deepwater trawl fisheries on seamounts and for seabirds taken in toothfish trawl and long-line fisheries, although mitigation measures have been implemented to reduce the bird catch (FAO, 2004b). The fragile and ancient coral “forests” found on seamounts are reduced to rubble by heavy trawl gear and consequently habitat for numerous other invertebrate species is lost. Improvements to trawl gear and monitoring may eventually allow the operation of deepwater pelagic trawls that avoid contact with bottom features, but in the short term the most effective mitigation measure to protect vulnerable and unique habitats is to close relevant areas to bottom trawling. The application of marine protected areas is controversial, especially in areas outside national jurisdiction, and the subject is under debate within the IUCN and CBD;6 notably, IUCN Recommendation 3099 calls for the protection of seamounts, deep-sea corals and other vulnerable deep-sea habitats from destructive fishing practices, including bottom trawling, on the high seas.7 Zoning the oceans into unfished marine reserves and areas with limited levels of fishing effort is one mechanism that might allow sustainable fisheries to be maintained within the diverse deep-sea ecosystems (Pauly et al., 2002). The inertia in developing and implementing international fisheries legislation has lead to NGOs calling for the designation of large-scale protected areas and for a moratorium on high seas bottom trawling until area management regimes can be implemented.8

5 http://www.ccamlr.org/pu/E/cds/intro.htm
8 http://www.savethehighseas.org/display.cfm?ID=136
A recent survey estimated that 47% of seamounts fall inside EEZs and 53% in international waters (Alder and Wood, 2004). Several countries have limited closures in place within their territorial waters. The Tasmanian seamounts reserve was voluntarily established in the Australian EEZ in 1996 and formally declared in 1999. Below 500 m the reserve has a protected area (IUCN management category 1a) and represents ~20% of the total seamounts in the local region. In the Tasman Sea, the Lord Howe Marine Park covers all the waters around Lord Howe Island down to 1800 m. Nineteen seamounts around New Zealand were closed to bottom trawling in 2000 (Clark et al., 2000). The closed seamounts were identified as being either representative of seamounts in their area or unique features in the EEZ. The faunal compositions of many seamounts are not known and selection of seamounts was based on geographical location and depth, rather than biodiversity. In the North Pacific the Bowie Seamount has a marine protected status that includes the conservation and protection of commercial and non-commercial fisheries. In Australia, the Department of Environment and Heritage have proposed establishing a network of large scale Marine Protected Areas (MPAs) in the SE marine region, that would cover 171 000 km² and close the major orange roughy fisheries (Buxton et al., 2006). Recent initiatives have lead to the establishment of a high seas Benthic Protected Area (BPA) in the Southern Indian Ocean, while representatives from the New Zealand fishing industry have proposed closing 31% of the seafloor (~1.2 m km²) in the New Zealand EEZ to bottom trawling, although the areas selected for closure need scientific review to ensure that the areas that are representative of marine environments in the EEZ.

7. SUMMARY
The expansion of fisheries into deepwaters, especially those in high seas outside coastal jurisdictions, has been the most significant development in world fisheries in recent years. Several of the important deepwater species are characterised by high longevity and late age at maturity; these species have ocean-wide distributions within a depth range and exhibit weak genetic differentiation within oceans. Several species, especially elasmobranchs taken as bycatch, are endangered; some teleost species in the NW Atlantic appear to have become endangered following 20 years of heavy exploitation. There is general agreement that many deepwater fishes are exploited beyond sustainable levels and that urgent action is required at the global level to reduce effort and to protect fragile ecosystems from the impact of bottom trawling. Difficulties in managing deepwater fishes are compounded by the high seas nature of many fisheries and the limited regimes available for management in international waters.

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Genetic resources for aquaculture: status and trends

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1. SUMMARY
Aquaculture, the farming of aquatic plants and animals, has grown consistently since 1970, when it provided only 3.9 percent of world fish supply. In 2004, global production of farmed fish (mainly crustaceans, molluscs and finfish) was over 45 million tonnes, comprising about 32 percent of total world fish supply, while the total production of farmed seaweeds for food and extraction of chemicals, was about 13.9 million t. Aquaculture also provides increasing proportions of the world’s supply of ornamental aquatic organisms. Over 90 percent of aquaculture takes place in developing countries, where it has high importance for poor people in terms of nutrition and livelihoods and where further responsible development of aquaculture, integrated with other natural resource use, has high potential for future growth. Based upon statistics submitted to FAO by its member States, about 84 percent of farmed fish production comes from Asia, with 67 percent coming from the Peoples’ Republic of China. However, aquaculture is increasing in importance in all developing regions and is expected to provide about 50 percent of world food fish supply within the next 20 years.

The future of aquaculture will depend in large measure upon the effective management of the genetic resources for farmed aquatic plants (PGR) and farmed fish (FiGR), as well as those for the organisms that provide their food and ecosystem services. Fish farms are agroecosystems and aquatic genetic resources for aquaculture on farms are part of agrobiodiversity. For example, microalgae and small invertebrates are mass cultured as live feeds for production of the early life history stages (“seed”) of farmed fish in hatcheries and natural feeds such as plankton are produced in fish farm waters. For some live feeds (e.g. the brine shrimp, *Artemia salina*) there is extensive information on genetic resources, but the genetic resources of most of the flora and fauna that support farmed fish production have been little explored.

The main difference between the status of most FiGR and aquatic PGR for aquaculture and all PGR and livestock (“farm animal”) genetic resources (FAnGR) for agriculture is that, with few exceptions, substantial domestication and genetic improvement of farmed aquatic species lag far behind the long history of purposeful breeding and genetic gains achieved for crops and livestock. This is now changing rapidly for some widely farmed aquatic species, such as tilapias, but much of the world’s production of seed for aquaculture and subsequent farm harvests remain documented mainly at the species level. Among the 80 species of livestock that are used for farming and ranching, over 6 000 different breeds have been recognized. The total number of aquatic animal species that have been farmed, experimentally or in actual production systems, is probably about 500, but the total number of farmed fish breeds has not yet been documented.

Many of the aquaculture statistics collected by governments and submitted to FAO are flawed; for example, by incomplete coverage of small-scale rural and
peri-urban aquaculture; by omission of data for some farmed aquatic species, such as freshwater macrophytes; by variable and incorrect nomenclature; and by aggregating and recording data by taxa higher than the species level. The relative importance of many genetic resources for aquaculture has still to be deduced in general terms from statistics that describe them as species, genera, families, commodity groups, and others “not elsewhere included (nei)”. For example, “aquatic plants nei” have become one of the largest contributors to production statistics for farmed aquatic plants. With few exceptions (e.g. catfish and striped bass), the contributions of fish hybrids, distinct strains, and other genetically altered forms are not yet recorded in most national statistics, and therefore cannot yet be accommodated in the statistics disseminated by FAO.

Information about genetic resources for aquaculture is not yet adequately covered by major global and regional databases and online information systems, including those currently provided by FAO and those that cover in detail the biology of aquatic organisms; e.g. FishBase. Moreover, there is a widespread need for greater standardization of correct nomenclature and terminology with respect to aquatic genetic resources. Progress is, however, underway in both these areas, with operators of databases and information systems for aquatic plants, crustaceans, molluscs and finfish now striving for greater collaboration and interoperability.

Major aquaculture publications and statistics reviewed from 1972 to 2004 suggest the following approximate ranges of numbers of farmable and potentially farmable aquatic organisms identified to species: microalgae, about 5 named as species, but with 16 genera also named; freshwater macrophytes, 5-8; marine macroalgae (seaweeds), 13-24; crustaceans, 26-79; molluscs, 20-74; other invertebrates, 4-7; finfish, 122-294; amphibians and reptiles, 3-11. Further exploration and documentation of the genetic resources of such large numbers of species - as wild and captive populations, geographical races, distinct farmed strains, hybrids and other genetically altered forms - will be a large task. However, the genetic resources for farmed aquatic plants could be covered under existing arrangements for terrestrial PGR and the most important FiGR for aquaculture could be prioritized; for example, by choosing initially the top 50 to 100 species that contribute most to farmed fish production, though with flexibility to include others that have clear potential importance and/or any wild and farmed FiGR that appear most threatened with extinction.

Consumer preferences are the main driver for farmers’ choices of which fish to farm. However, most of the world’s aquaculture and culture-based fisheries production is based on seed produced from broodstock populations by the operators of fish hatcheries. Public and private seed producers, their breeding programmes and related research determine largely which types of seed are available for purchase by farmers, for subsequent growout to marketable size. Fish farms range in size from small-scale/backyard to large scale corporate ventures. Vertically integrated aquaculture, similar to broiler chicken production, is also expanding. Most aquaculture is undergoing intensification to boost production per unit area or volume of farm waters. This requires the development of strains, hybrids and other genetically altered forms that are tailored to intensive farming, especially with respect to commercial traits such as good feed conversion, disease resistance, fillet yield, colour, flavour, etc.

Because of the short history of domestication, breeding programmes and related research for most farmed aquatic organisms, the free-living populations of their wild and feral relatives and of other potentially farmable aquatic species have high importance as genetic resources. Many of these free-living populations, especially in freshwaters, are among the world’s most seriously threatened biodiversity; for example, the wild genetic resources of farmed carps and tilapias. Moreover in aquaculture, as in agriculture, most private sector seed producers and farmers keep only the most profitable farmed species and types, leaving others under threat of extinction. The use
in aquaculture production and related research of alien species and of genetically altered forms (e.g. distinct strains, hybrids, polyploids, transgenes etc., whether developed from alien and/or indigenous species) is certain to increase. This will require more effective biosafety and biosecurity procedures than have been implemented to date, particularly with respect to thorough appraisal of the impacts of escapes and releases of farmed aquatic organisms before granting approvals for introductions and transfers, as well as strictly enforced quarantine.

These trends indicate an urgent need for better management – meaning fully integrated use and conservation – of aquatic genetic resources for aquaculture: in situ/in vivo, as free-living, wild and feral populations; in situ/in vivo, as captive populations on-farm; ex situ/in vitro, as collections of cryopreserved sperm, embryos and other tissues/DNA; and ex situ/in vivo as aquarium and research populations. This will require increased investment in the management of FiGR and aquatic PGR, commensurate with their high and growing contributions to world food security, Keeping representative, free-living wild populations of farmed fish species undisturbed in their natural habitats and off-limits to aquaculture and to contact with farmed fish, has operational and opportunity costs. Therefore, unless there is equitable sharing of costs and benefits among the stewards and potential users of such aquatic genetic resources for aquaculture, the conservation element in their management will not be achieved. Establishing and maintaining ex situ, in vivo and/or in vitro, fish gene banks is also expensive and will require public and private sector investment and partnerships. Attempts by the private sector to acquire intellectual property rights on genetically altered fish and related biotechnological processes in aquaculture have so far been limited, compared to the situation in plant breeding. It is unlikely that attempts to enforce proprietary rights on genetically altered fish will prosper in the near future. Rather, as public and private fish breeding programmes develop, returns to fish breeders will likely come from purchased access to pedigreed fish populations and eventually to pedigree individuals, as for livestock and pet animals. However, private sector research, especially for the development of biotechnological products and processes, is bound to increase in aquaculture, following the trends in agriculture.

The following strategic directions are suggested for improving the management of genetic sources for aquaculture: increased investment; management (i.e. fully integrated use and conservation) as part of agrobiodiversity; improved information systems; conservation in changing ecosystems; reconciliation of aquaculture with nature conservation; progressive linking of the management of aquatic PGR and FiGR with that for terrestrial PGR and FAnGR; and exploration of the application of an interactive governance approach, with assessments of the governability of aquatic genetic resources.

2. INTRODUCTION

Aquaculture is the farming of aquatic plants and animals. It comprises the mass production, usually in hatcheries, of “seed” (eggs, larvae, postlarvae, fry, fingerlings, juveniles etc.) of farmed aquatic organisms, and the subsequent growout of that seed to marketable size in aquatic farms or its release for culture-based fisheries (CBF) (e.g. see Bartley and Leber, 2004; Caddy and Defeo, 2004). Hatchery operations for CBF are generally considered part of aquaculture. The FAO Code of Conduct for Responsible Fisheries (FAO, 1995) and its guidelines for aquaculture development (FAO, 1997) refer throughout to “aquaculture, including culture-based fisheries”. Seed is produced mainly from captive breeding populations. However, for the minority of farmed aquatic species where mass production of seed in captivity is not yet technically possible, or where its collection from wild populations still makes economic sense, wild seed or young adults are obtained from capture fisheries and then grown to marketable size in captivity. This can be termed capture-based aquaculture (e.g. Ottolenghi et al., 2004).
This review is concerned mainly with the genetic resources of fish, meaning finfish and aquatic invertebrates (principally crustaceans and molluscs) that are farmed or potentially farmable. The genetic resources for CBF, as well as their genetic impacts on wild populations, are not considered here. Most farmed aquatic plants and animals are used for human consumption as food but some are farmed for other purposes; e.g. for extraction of industrial chemicals (seaweeds), as ornamental species (aquatic plants, invertebrates, finfish, amphibians and reptiles), for sport fisheries (finfish) and for cosmetic, jewelry, and medicinal products (molluscs, seahorses etc.). It is implicit in this review that policy and other provisions made for the genetic resources of aquatic organisms farmed as human food should apply also to those of aquatic organisms that are farmed for other purposes. Genetic resources for farmed aquatic plants are covered briefly here, emphasizing macroalgae (seaweeds) farmed for human food or for extraction of chemicals. All genetic resources for farmed aquatic plants are called PGR.

By convention, all fish genetic resources for aquaculture and capture fisheries are now termed FiGR. FAO aquaculture statistics include farmed macroalgae within a general definition of "fish", but their genetic resources are PGR, not FiGR. Farmed aquatic amphibians and reptiles also figure in FAO and some other farmed fish statistics, but can be considered as livestock ("farm animal") genetic resources (FAnGR), thereby restricting the use of the term FiGR for farmed aquatic vertebrates to finfish alone. Similarly, the farming of aquatic birds and mammals is not considered part of aquaculture, and their genetic resources are regarded as FAnGR, not FiGR. Farmed amphibians and aquatic reptiles are mentioned here only insofar as they are included in FAO aquaculture statistics and major texts.

This review builds upon recent publications that address conservation and use of aquatic genetic resources (e.g. Pullin et al., 1999; Pullin, 2000, 2006b; Science Council Secretariat, 2005). The importance of aquaculture, its rapid growth and dynamic nature are summarized, with overviews of the main categories of genetic resources for aquaculture; i.e., for feeds and ecosystem services, aquatic plants and fish. Discussions follow on factors that affect the status of and trends in genetic resources for aquaculture: choosing what to farm; information and nomenclature; threats; management, defined as fully integrated use and conservation; and the sharing benefits and costs, including ownership and use issues. No order of priority is implied here. The review concludes by identifying some strategic directions for improving the management (i.e., the fully integrated use and conservation) of genetic resources for aquaculture.

3. THE GROWING IMPORTANCE OF AQUACULTURE
FAO is the source of all aquaculture statistics quoted here, unless otherwise stated. FAO began to publish statistics in 1950 but, up to 1984, aquaculture statistics were combined with those for fish catches. Despite their subsequent separate status and increasing importance, world aquaculture statistics are still beset with uncertainties. There is a widespread need to improve collection of data from small-scale, rural aquatic farms, especially in developing countries. The world’s small-scale rural and peri-urban aquaculture production, as well as its value and importance in household food security and provision of incomes and employment are probably substantially under-recorded in many national statistics. Moreover the real, as opposed to perceived, contributions of many CBF to world fish supply are poorly known and will remain so unless data for their seed production and harvests are adequately disaggregated from those for growout on farms and capture of wild fish. Uncertainties concerning the current contributions and future potential of CBF have been mentioned by many authors (e.g. Lorenzen et al., 2001; Leber et al., 2004). There is also a need to analyse trends in aquaculture both with and without inclusion of the statistics reported by the Peoples’ Republic of China (PRC) (e.g. New, 2003).
Despite these uncertainties, the present contributions of aquaculture to world food security and its future potentials are well recognized. Aquaculture has large potential for further growth, not only in the countries where it is well-established but also in many of those where it is relatively new, including sub-Saharan Africa and Latin America. Governments in all developing regions have framed and begun to implement policies that place reliance on expansion and intensification of aquaculture for sustaining and increasing their fish supply (e.g. see Brugère and Ridler, 2004).

In 2002, the status and future prospects of aquaculture were described as follows in a background paper for the first meeting of the FAO Sub-Committee on Aquaculture (FAO, 2002):

“Aquaculture is an important domestic provider of much needed, high quality, animal protein, generally at prices affordable to the poorer segments of society. It is also a valuable provider of employment, cash income, and foreign exchange, with developing countries contributing over 90 percent of the total global production. When integrated carefully, aquaculture also provides low-risk entry points for rural development and has diverse applications in both inland and coastal areas.”

Annual rates of increase for aquaculture production and value have varied greatly with species and farming systems but, since the 1970s, almost all have been higher than those for other food production sectors and remain so. For example, shrimp farming in the late 1970s grew at 24 percent per year and FAO (2002) described its 6 percent average annual growth rate in the 1990s as “modest”. Farmed fish currently provide about 32 percent of world food fish supply, compared to about 3.9 percent in 1970 and their contributions are widely expected to grow to about 50 percent, probably within the next 20 years. According to McHugh (2003), most of the world’s production or macroalgae for human food and for extraction of chemicals (hydrocolloids) is derived from aquaculture. For 2004, FAO statistics indicate total world production of 13.9 million tonnes of farmed aquatic plants, worth about $6.8 billion. Aquaculture is also an increasingly important source of supply for ornamental freshwater and marine tropical fish, in developed and developing countries. Information on ornamental plants and animals is widely available through global databases (e.g. for marine fish and invertebrates, see www.unep-wcmc.org).

A nutrition transition, from diverse, traditional fish-, fruit- and vegetable-rich diets to fat-, sugar- and alcohol-rich diets, is underway in the developing world and is causing rapid growth of diet-related, chronic diseases (ischemic heart disease, diabetes, obesity, hypertension, stroke, and certain cancers), with high consequential costs. In 1995, these diseases accounted for 41.6 percent of all deaths and 22.5 percent of all hospital expenses in the PRC, equivalent in total to 2.1 percent of gross domestic product (GDP), while for Sri Lanka the corresponding figures were 18.3 percent, 16.7 percent and 0.3 percent of GDP (Popkin et al., 2001). Gillespie and Haddad (2001) reviewed the “double burden” of malnutrition: undernutrition and overnutrition from overconsumption of unhealthy foods. Farmed fish will be increasingly important contributors in efforts to solve these problems, especially as they can provide substantial nutritional and livelihood benefits to the poor (e.g. ADB 2005a; FAO/NACA-STREAM 2005). For many developing countries, aquaculture is the main hope for sustaining and increasing contributions of affordable fish and fish products to healthy diets. Fish provide their consumers with animal protein, health promoting lipids and essential vitamins and minerals and are particularly important in human nutrition as sources of the omega-3 fatty acids necessary for brain development in the human foetus and its functioning throughout life (e.g. Elvevoll and James, 2000; Anon., 2006).

Aquaculture is often categorized according to the feeds available to farmed fish. In extensive aquaculture, fish depend entirely on the natural productivity of farm waters,
supplying natural feeds: plankton, detritus, vegetation etc. In semi-intensive aquaculture, relatively cheap supplementary feeds are given, and the production of natural food in farm waters is sometimes artificially increased by fertilization. In intensive aquaculture, farmed fish are entirely dependent upon provision of nutritionally complete feeds, which typically account for about 65 percent of the total variable costs of production. Intensification, through maximizing use of pond fertilizers and supplemental feeds to intensive feedlot systems, is now a major trend for most forms of aquaculture. This boosts production per unit area or volume of farm waters, but makes large ecological footprints beyond farming areas. The main exceptions to this are seaweed farming and most farming of bivalve molluscs, which remain largely extensive aquaculture operations, involving minimal husbandry from seed to harvest.

Table 1 summarizes the most recent aquaculture production and value statistics (2004), by the top 10 leading countries and the rest of the world, for fish farmed for human food. From these data, Asian countries accounted for 84 percent of world aquaculture production in 2004, with the PRC alone accounting for 67 percent. Note the higher values accorded to aquaculture produce in the more developed countries.

### 4. GENETIC RESOURCES FOR AQUACULTURE

#### 4.1 Feeds and ecosystem services

All sources of human food production, including aquaculture, are interconnected as a global food web. The genetic resources for the cereal crops and other plants that provide ingredients for the feeds given to farmed fish are genetic resources for aquaculture. Similarly, the genetic resources for the low value/trash fish (LV/TF) and industrial fisheries that provide fish, fishmeal and fish oils for feeding farmed fish and livestock are genetic resources for both aquaculture and livestock production. However, Tacon et al., (2006), citing FAO (2005), pointed out that only 18.2 percent of global fishmeal production and 45 percent of fish oil production is currently attributable to named species. This means that many of the FiGR for fishmeal are fish oil production are undocumented, even at species level. From a world food security perspective, it is important to note that aquaculture production which remains based upon substantial use of wild caught fish, fishmeal and fish oil, cannot be claimed as a net gain in fish supply or as a net contribution to filling the gap in fish supply caused by declining capture fisheries. Tacon et al., (2006) estimated that in 2003 the “aquaculture sector”
Genetic resources for aquaculture: status and trends

Consumed as feeds the equivalent of 20 to 25 million tonnes captured fish, as live weight equivalents, in order to produce 30 million tonnes of farmed finfish and crustaceans. They identified the following groups of farmed fish as net consumers or producers of fish: net consumers – river eels, marine fish and shrimps, salmon and trout; net producers – carp, catfish, freshwater crustaceans, milkfish and tilapia.

Production of fish seed in aquaculture and for CBF often involves protein-, essential lipid- and micronutrient-rich starter fish feeds; supplied in fine particulate form or as live food organisms, cultured or collected specifically for this purpose; e.g. bacteria, microalgae, rotifers, crustaceans and molluscan larvae. The genetic resources for organisms that are used to produce these feeds and for live food organisms used in aquaculture are also genetic resources for aquaculture. The status and diversity of some of the latter are well-documented; for example, there are interlinked collections and information sources for cultured bacteria and microalgae and a reference centre for the brine shrimp *Artemia salina* and the rotifer *Brachionus plicatilis* (www.aquaculture.ugent.be).

Many other microbial, plant and animal species provide farmed fish with food and feed ingredients and with a wide range of ecosystem products and services, including oxygen, shelter, spawning substrates and waste processing. Their genetic resources are essential for the future of aquaculture, being broadly analogous to the genetic resources for organisms that contribute organic fertilizers for the production of crops and fodders for livestock. Indeed, all species that provide feeds and ecosystem services to aquaculture are part of agrobiodiversity when found on-farm; i.e., in agroecosystems. These supportive genetic resources for aquaculture merit much wider recognition and documentation, and above all more effective management, than they have received to date.

4.2 Farmed aquatic plants

Farmed aquatic plants comprise green microalgae (e.g. Chlorella); blue-green algae, more properly termed cyanobacteria (e.g. Spirulina); macroalgae (brown, green and red seaweeds); and freshwater macrophytes (e.g. floating species, such as azolla and duckweeds, and emergent species such as lotus, water chestnut and water spinach). Table 2 gives numbers of farmed aquatic plants identified to species in some major aquaculture publications.

Farmed microalgae are not well covered in most aquaculture literature, except as live feeds for fish hatchery operations. FAO statistics give production of farmed Spirulina in 2004 as 41,750 tonnes. *Chlorella vulgaris* is listed, but with zero production recorded. Stickney (2000) mentioned 16 farmed microalgal genera.

<table>
<thead>
<tr>
<th>Farmed aquatic plants</th>
<th>Numbers of species</th>
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<tr>
<td></td>
<td>A</td>
</tr>
<tr>
<td>Microalgae</td>
<td>5</td>
</tr>
<tr>
<td>Freshwater macrophytes</td>
<td>8</td>
</tr>
<tr>
<td>Marine macroalgae (seaweeds)</td>
<td>15</td>
</tr>
<tr>
<td>TOTAL</td>
<td>28</td>
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Source: A: Bardach et al. (1972); B: Pillay (1990); C: Stickney (2000); D: FAO aquaculture statistics (2004)
Despite their high importance as human food, as fodders and fertilizers in agriculture and as components of waste treatment systems (e.g. Edwards, 1980; Van Hove, 1989; Kanungo et al., 2001), farmed freshwater macrophytes are not well covered in mainstream aquaculture literature and FAO aquaculture statistics. Some freshwater macrophytes – for example, water spinach (*Ipomea aquatica*) are major crops, but information on their production and their genetic resources is not easily obtained, either from agriculture or aquaculture sources. Conversely, the wetland and deepwater rice, which are aquatic macrophytes, are well covered by mainstream crop genetics literature. Most of the available information on other freshwater macrophytes concerns control of nuisance species; for examples, see the Journal of Aquatic Plant Management; http://www.apms.org/japm/japmindex.htm. However, a new “forum” about peri-urban farming of freshwater macrophytes and fish is being established, based upon examples in Southeast Asian cities (contact: W. Leschen; wl2@stir.ac.uk).

FAO statistics for farmed aquatic plants focus on marine macroalgae (seaweeds) and are included with farmed fish statistics. They name only eight macroalgal species and group others together within seven genera and/or as higher taxa. The major contributors to world farmed seaweed production that are identified to species are *Laminaria japonica*, *Porphyra tenera*, and *Eucheuma cottonii*. Large contributions are said to come from “aquatic plants nei” (i.e. not elsewhere included), which are assumed to be macroalgae. Production of these aquatic plants nei has tended to increase, mainly because of the larger quantities reported from the PRC since 1998 (1 946 980 tonnes) as compared to 1997 (461 675 tonnes). Prior to 1998, production of farmed seaweeds in the PRC was reported on a live (wet) wet basis, whereas from 1998 it was recorded first as dry weight and then reported after applying conversion factors (A. Lowther, personal communication). Figure 1 shows the trends in production of the four major farmed species, plus aquatic plants nei, from 1985 to 2004.

McHugh (2003) forecast limited scope for expansion of seaweed farming as follows: to supply agar, limited; to supply alginites (typically from *Laminaria japonica*), about 2-3 percent per year; to supply carrageenan, about 5 percent per year; and as human food, highly variable prospects, dependent upon promotional efforts. However, seaweed farming undoubtedly has potential to improve the lives of some poor and marginalized
coastal communities, especially in the tropics. For example, in the Philippines Autonomous Region in Muslim Mindanao, some poor coastal communities in the farm seaweed as contract growers, for exporters of seaweed products. In 2004, this region produced 472,514 tonnes of farmed seaweed; over 50 percent of the Philippine national total of that used for exportable seaweed products (Unson, 2006). Against the many actual and potential benefits of seaweed farming, there is serious cause for concern when alien macroalgal species are introduced for aquaculture to new coastal locations without through prior appraisal of their possible ecological impacts.

More detailed coverage of production and value data for farmed aquatic plants, with authoritative and correct names at species level, is an essential prerequisite for monitoring the status of and trends in their genetic resources. This merits high priority, not only for the major commercial species groups but also for those that are of high importance as contributors to the food and livelihood opportunities of poor communities; e.g. *Caulerpa* spp. in tropical Asia. The database www.algaebase.org is a good source of information on correct taxonomy and nomenclature of algae and could be supplemented to give information on the genetic resources of farmed algae. At present, however, most information about these PGR is scattered and is to be found mainly in the major phycological journals and occasionally in those that cover aquaculture in general (e.g. Cheney 1999). It could be collected and made accessible through existing arrangements for terrestrial PGR, given additional investments.

### 4.3 FARMED FISH

FiGR for aquaculture can categorized in a wide variety of ways: by taxonomy and genetic terminology (e.g. allele, selected strain, hybrid, artificial polyploid, transgene, species, genus, family, order, commodity group etc.); by location (area of production; natural and introduced geographic ranges; by free-living and/or farm environments, including migratory habits (brackishwater/diadromous; freshwater; marine) and production systems (cages, pens, ponds, raceways, recirculating systems, tanks, etc.); by relative current worth (production tonnages, monetary values, nutritional importance, poverty alleviation through livelihood provision and diversification, sociocultural value; sport and recreational value etc. However, the main basis for categorization of FiGR for aquaculture is their actual and potential use, as indicated by aquaculture statistics and research findings. Table 3 gives numbers of farmed aquatic animals identified to species in some major aquaculture publications.

#### TABLE 3
Approximate numbers of farmed aquatic animals that are named as species in some major aquaculture publications, including those having potential for farming

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<thead>
<tr>
<th>Farmed aquatic animals</th>
<th>Numbers of species</th>
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<td></td>
<td>A</td>
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<tr>
<td><strong>Invertebrates</strong></td>
<td></td>
</tr>
<tr>
<td>Crustaceans</td>
<td>79</td>
</tr>
<tr>
<td>Molluscs</td>
<td>61</td>
</tr>
<tr>
<td>Others (mainly echinoderms)</td>
<td>-</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td>140</td>
</tr>
<tr>
<td><strong>Vertebrates</strong></td>
<td></td>
</tr>
<tr>
<td>Finfish</td>
<td>294</td>
</tr>
<tr>
<td>Amphibians and reptiles</td>
<td>6</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td>300</td>
</tr>
<tr>
<td><strong>TOTAL</strong></td>
<td>440</td>
</tr>
</tbody>
</table>

Sources: A: Bardach et al. (1972); B: Pillay (1990); C: Nash (1993); Nash and Novotny (1995); D: Stickney (2000); E: FAO aquaculture statistics (2004)
FAO aquaculture statistics retain data entry lines for species and higher taxa for which zero production and value are recorded. For some, production has been zero for decades. This means that FAO’s aquaculture statistics are records of all historical use of these species and higher taxa, not just records of recent and current farming. Figure 2 shows the production of farmed fish by major groups (crustaceans, finfish and molluscs) from 1985-2004, with production of other farmed aquatic invertebrates and of farmed aquatic amphibians and reptiles seen as very small by comparison.

As Bartley et al., (2001) have shown, interspecific fish hybrids are used in aquaculture, but their contributions to production go largely unrecognized and, with very few exceptions (e.g. hybrid catfish [Clarias gariepinus x Clarias macrocephalus] and hybrid striped bass [Morone chrysops x Morone saxatilis]), are not yet captured adequately in national or FAO statistics. The data from member countries, upon which FAO statistics are based, is given only at species level or at higher taxa comprising unspecified numbers of species; for example, genus + “spp.” and “not elsewhere included”. There is no information concerning any taxon below species level.

FAO statistics can be analysed in various ways to attempt to prioritize farmed aquatic species. Contributions not only to aquaculture production and value but also to availability of produce that is affordable by poor consumers would probably be the most equitable and best broad measure. Such prioritization would, however, be a lengthy exercise and is not attempted here. A good general idea of the approximate numbers of important farmed fish can be gained from recent analyses. For example, New (2003) lists the following numbers of clearly important species: 8 crustaceans; 10 molluscs; and 26 finfish (13 freshwater, 7 diadromous and 6 marine). This gives a total of 44 most important species, but more flexibility and inclusiveness are needed to prioritize FiGR for aquaculture because some species are of special importance to only one or a few countries.

The relative national and international importance of a farmed aquatic species can change rapidly; for example, farmed Nile tilapia (Oreochromis niloticus) production has shown extraordinarily rapid growth in recent years (Figure 3), though a substantial proportion of what is currently recorded as production of farmed Nile tilapia is
probably of tilapia hybrids having this species as one of the parents. The Pacific white shrimp (Litopenaeus vannamei) has rapidly become the main species of farmed penaeid shrimp. The data shown in Figure 4 are probably underestimates of its increasing contributions, because some countries reporting to FAO take time to adjust their reporting by species as the proportions of farmed species change. L. vannamei probably now accounts for over 80 percent of farmed penaeid shrimp production in Asia.

The world’s FAnGR for livestock farming and ranching comprise about 80 species, of which 14 contribute most to world production and within which over 6000 breeds have been recognized, whereas the world’s FiGR for aquaculture probably comprise about 500 species that have recorded as having been farmed to some extent (including experimentally) at some time (Pullin, 2006b; Science Council Secretariat, 2005). FishBase (www.fishbase.org) has listed 344 species of farmed finfish. However, data currently available at species level in aquaculture statistics and aquaculture research literature suggest to this author that substantial coverage of FiGR for aquaculture could be achieved by prioritizing 50 to 100 species of farmed and potentially farmable fish, taking into consideration their international and national importance as well as their status, especially where they are threatened (see 7. below).

All major livestock species are considered fully domesticated. Their few remaining wild relatives are of low importance for future breeding programmes, and there are few new potential candidate species for farming. Most farmed fish species are not yet fully domesticated. Their wild relatives are of high importance for breeding programmes and related research, and there are many (possibly hundreds) of new potential candidate species for aquaculture. Balon (2004) argued that only the common carp (Cyprinus carpio) as a farmed food fish and as koi ornamental carp, the goldfish (Carassius auratus) and a few other ornamental species can be called true domesticates, with other farmed fish (including Chinese and Indian carp, catfish, salmon, sturgeon and trout species) qualifying only as “exploited captives”, apart from their few colour variants, such as albino strains, that can be termed “domesticated”. There is good evidence to support this view. For example, the diversity and stability of goldfish (Carassius auratus) breeds are comparable to those for dog breeds (e.g. Zhen, 1988), but most
farmed fish strains and hybrids look alike and the consumer of farmed fish and farmed fish products does not yet have breed-specific choices, comparable to those available for many livestock products. At present, the world’s farmed fish are represented by relatively few well-documented, distinct and stable breeds.

Prior to the big expansion of application of genetics in aquaculture that began in the late 1980s, development of and documentation about distinct and stable breeds and hybrids of farmed fish were poor. Even by the 1990s, fish breeds and hybrids had not been developed for particular farm environments and farming methods and for most farmed aquatic species, particularly in the developing countries, well-documented FinGR of known provenance were simply not available. This meant that the products of any well-reputed genetic improvement research were almost certain to enjoy high demand for use in a wide range of farming systems. GIFT and GIFT-derived Nile tilapia are a good example. Having been bred from initial research trials in a wide range of farm environments from ricefields to ponds and cages (Eknath et al., 1993), GIFT and GIFT-derived Nile tilapia have been farmed in most tilapia farming systems and, in view of their broad genetic base, have become the main basis for national tilapia breeding programs in several countries (ADB, 2005b).

Parallel to the intensification of aquaculture, there is an ongoing quest to push many farmed aquatic species towards omnivory and acceptance of least-cost formulated feeds, irrespective of their natural feeding habits (Pullin, 2006a). Many farmed fish, especially marine species, are naturally carnivorous but are being constrained to accept feeds containing as much plant and microbial protein as is biologically possible, as well as a wide range of rendered livestock and other waste products. Conversely, many widely farmed and naturally herbivorous and omnivorous fish species (such as grass carp and Nile tilapia) are being farmed more and more intensively, using feeds containing fishmeal, rendered livestock products etc. In general, these trends require the development of fish strains, hybrids and other genetically altered forms that perform well in intensive farming systems, that show good feed conversion on low cost feeds, that yield attractive and well-flavoured products, and that enjoy high survival
and growth performance in adverse environments; for example, cold- and saline-tolerant tilapias. Breeding programmes and related research that compare these and other commercial traits among different farmed strains, hybrids and other genetically altered forms are therefore increasing (e.g. Rutten et al., 2004a, 2004b) and will draw upon FiGR from farmed, wild and feral populations, including those established in adverse environments. Costa-Pierce (2003) recognized the importance of feral tilapia populations and recommended establishment of a registry using genetic markers. Over the past 30 years, fish breeding programmes and related research have been undertaken largely by public institutions and organizations, but will be increasingly pursued by public-private partnerships or by the private sector alone.

5. CHOOSING WHAT TO FARM
Genetic improvement of farmed fish lags far behind genetic improvement of crops and livestock but is taking similar approaches. Crop breeding and related research are increasingly driven by market assessments of demand for certain types of seed, with the development and importance of different genetic resources (varieties, hybrids, etc.) determined mostly by demand-led technical change, rather than supply-led proposals from scientists (P. Pardey, personal communication). The same trends are likely to develop in fish breeding.

Fish consumers determine the demand for different types of farmed fish at any given time, while aquaculture science works to develop and to introduce new options. Most fish consumers are, however, unaware of the existence and importance of FiGR. They usually buy, or receive (for example, in disaster relief operations) aquatic produce that they know only by common names. Their categorization and choices of produce usually approximate to species level, though they often know the names of the places of production of farmed fish (e.g. Scottish salmon in the United Kingdom; Batangas Province tilapia in the Philippines) and seek produce from a named location, based on their previous experience of buying it or on perceptions about its quality. The naming of places of production in fish markets, as in fish restaurant menus, is often a marketing ploy and does not usually provide reliable information about the genetic identity of produce. For example, some of the salmon farmed in Scotland and other countries were bred in Norway, and many farmed salmon look alike irrespective of origin and breeding history. In developing countries, there is rarely any independent certification that fish in the market place bearing the name of an area or farm of origin all came from there.

In many countries, though primarily at present in the developed world, consumers’ choices of farmed fish are being made increasingly on ethical grounds. Ethics and responsibility in aquaculture have been reviewed by Kaiser (2002). For fish consumers, the main factors are whether farmed fish are treated humanely and whether they are produced in environment- and biodiversity-friendly farming systems; considering not only the obvious impacts of effluents from fish farms, abuse of antibiotics, etc., but also the choice of fish with feeding requirements – preferably herbivorous/omnivorous – that will not exacerbate pressures on capture fisheries that are already overexploited. Public perceptions of genetic modification of food species are also a major factor in ethically-based choices of what to eat, irrespective of considerations of biosafety and food safety. All such ethical considerations are being applied to farmed fish, particularly as organically farmed fish are becoming new entrants to organic agriculture (www.ifoam.org). Fish welfare issues, including those of farmed fish have been reviewed by Huntingford et al. (2006).

In most aquaculture, as in most agriculture, seed production and growout are separate enterprises, in different hands. Also in aquaculture, as throughout agriculture, seed producers’ and farmers’ choices of which aquatic organisms to farm are
determined by market demand, profitability, and technical feasibility. Assessments of all of these imply risk assessment and management, and these in turn require information as well as adequate knowledge and skills. Seed producers and farmers base their choices of fish upon their own experience and/or external advice concerning a wide range of commercial traits: e.g. for seed producers, fecundity of and egg quality from broodstock, and survival, growth rate and disease-resistance of seed; for farmers, survival, growth rate, feed conversion, disease resistance, dressing weight, color, flavour etc. Many farmers, especially small-scale farmers in developing countries, have to make choices about what to farm while lacking adequate science-based information and independent advice on the genetic diversity that is available and on the performance of different species, hybrids and strains. The links in the “chain of choice” concerning what to farm are at their strongest in modern, vertically integrated aquaculture and agriculture, where research, breeding, seed production, contract growing, processing and marketing are all or mostly undertaken within the same organization – usually a large food company which also manufactures feeds and supplies technical support services. Some forms of aquaculture, such as intensive farming of Nile tilapia, already resemble vertically integrated poultry farming though, like chickens, tilapia can also be farmed in a wide range of systems from free range through backyard feedlot to small, medium and large scale commercial farms (e.g. see Young and Muir, 2002).

Although choices about which fish are farmed are primarily consumer-driven, many other actors, including researchers, breeders, and fish processors, also influence these choices. Decision-making along this chain is a research area that has been little explored, but it is probable that some of its links are weak or even disconnected. Most fish consumers, and indeed fish farmers, feel that they know what need, while researchers, breeders and seed producers tend to promote their new ideas and products, often with strong political and commercial backing. Sometimes this results in large benefits to farmers and consumers, sometimes not. A good positive example was the development of new technology for the farming of genetically improved farmed tilapia (GIFT) (ADB, 2005b). However, interactions among aquaculture scientists, seed producers, farmers and fish consumers are often weak. Globalization is increasing the remoteness of some fish farmers from their markets. For others who remain closer to their markets, consumer demand and profit margins clearly dictate the choice of what to farm. An important recent example can be seen in the switch made by carp farmers in Andra Pradesh, India, from following long-established, scientist-derived polyculture formulae, that required stocking six (three indigenous and three alien) carp species in all ponds, to a much simpler system of stocking just two indigenous carp species, resulting in greater yields and profits (Nandeesha, 2001). This worked because of the high price of one of these species (Catla catla) and the opportunistic feeding behaviour of the other (Labeo rohita). The theoretical basis of multispecies polyculture – different species occupying separate feeding niches (benthos, detritus, phytoplankton, zooplankton, etc.) – tends to break down as aquaculture is intensified.

The other main actors whose activities influence current and future choices of what to farm, as well as where to farm it, are the conservationists at all levels (international, national and local/community) who recognize the need to conserve not only the genetic resources of farmed aquatic organisms, but also those of their wild relatives, of farmed types for which production has been discontinued, and of potential new candidate species for aquaculture. The overall goal here is to maximize options for future availability and use of FiGR and aquatic PGR. In agriculture, conservation of the wild relatives of farmed plants and animals and of traditional and rare varieties and breeds seems to be generally of less importance than it is in aquaculture. Moreover in agriculture new candidate species for farming are few, whereas in aquaculture there are probably hundreds. For aquaculture therefore, with its limited history of documentation and development of genetic resources, there is a strong case for
assuming that all distinct wild, feral and farmed populations of farmed and potentially farmable aquatic species are potential sources of unique and useful genetic material for aquaculture. However, choices also have to be made among this vast array of genetic resources. Those choices will again be largely influenced by the current choices of consumers as well as the opinions and foresight of researchers and breeders and other actors in the chain.

6. INFORMATION AND NOMENCLATURE

6.1 Crossing communication barriers

Broadly speaking, aquaculture researchers and most fish breeders talk the language of science and understand genetic terminology, whereas many seed producers and farmers and almost all of the general public do not. Inevitably, there is a mismatch between how scientists document genetic diversity in aquaculture and how most seed producers, farmers and consumers perceive, categorize and name farmed fish. The same applies to the conservation of wild populations, for some of which there is a rich folk taxonomy in local languages (e.g. see May, 2005) as well as a rapidly increasing reliance on molecular genetic data (e.g. see Hedrik, 2004).

Common names are the most obvious way through this barrier. FAO uses common names extensively in its provision of fisheries information, including aquaculture statistics. FishBase (www.fishbase.org) provides authoritative and correct nomenclature at species level for finfish, with user entry possible through the scientific names of fish and through their common names in over 200 languages. However, many of the common names listed by FAO, FishBase and others are highly contrived, for the simple purpose of just assigning a name other than a scientific name, which can be daunting to lay users. Therefore many so-called common names are not actually in common use. For example, the tilapia *Sarotherodon galilaeus* is listed by FAO and FishBase as the “mango tilapia”, with FishBase suggesting the USA as the source of this common name. This is a beautiful name, but this author has never heard it used anywhere.

More serious problems with nomenclature can occur when the collectors and compilers of aquaculture statistics fail to keep abreast of changes in the scientific nomenclature of farmed aquatic organisms. Taxonomists are constantly revising nomenclature and often disagree about the status of species, which means that at any given time some diversity in nomenclature is inevitable. Recognizing this, the world’s taxonomic databases and information systems increasingly allow not only for entry through common and scientific names but also provide coverage of synonyms and common misspellings of the latter to assist users to find the information that they seek, and also to consider correcting their nomenclature thereafter. FishBase has long provided such coverage for finfish and it is also available in global databases such as the Catalogue of Life (www.sp2000.org) and Namebank (www.ubio.org). The phylogenetics database Deepfin (www.deepfin.org) links finfish systematicists as a research coordination network and is a useful source for nomenclatural changes.

Overall, the goal for all concerned with management of information about genetic resources for aquaculture must be to call all farmed aquatic species, as far as is possible, by their correct scientific names. For some widely farmed fish this is not yet done rapidly. For example, the mrigal, an important farmed carp species, is not yet widely listed under its correct name *Cirrhinus cirrhosus*. Where taxonomic revision has involved splitting or lumping species, some statisticians persist in using old and incorrect names which fail to indicate the importance of what have come to be recognized as the same species or as separate individual species. A good example of the former is the widely farmed silver barb, an Asian carp, now properly called *Barbonymus gonionotus*. It was formerly called *Puntius gonionotus* or *Barbodes gonionotus*, names which are still found in some statistics and research papers. However, the main problem here is that
some aquaculture statistics still refer erroneously to and list separate data for another species, the Java barb or Puntius javanicus, all populations of which are now known to be Barbonymus gonionotus.

As a further example of the need to check nomenclature, even in international centres of excellence for research and development, in 1999, a FishBase team checked the correctness of all of the scientific names of plants and animals used by the 16 centres of the Consultative Group on International Agricultural Research (CGIAR), including those entered in its System-wide Information Network for Genetic Resources (SINGER) (ICLARM, 2000). The names used by the CGIAR centres and SINGER were compared with the most authoritative sources available; e.g. the Germplasm Resources Information Network (GRIN) and Species 2000. The results were revealing: for example, 3,183 SINGER names did not match valid names or synonyms in GRIN; 400 names used in the SINGER matched synonyms or known misspellings in Species 2000; and 960 SINGER names had no matches in GRIN or Species 2000. It is vital to check all names entered into statistical and other databases that will be used for making policy and decisions about use and conservation of FiGR. Only then will all synonymies and common misspellings be revealed and understood and databases that use scientific names as entry points be fully linkable. Standardized and correct nomenclature at species and interspecific hybrid levels is the first step, before venturing into intraspecific taxa and molecular genetic terminology, which must also be correct and, as far as is possible, standardized.

6.2 Information sources, gaps and future needs

Substantial information about FiGR for aquaculture has been and will continue to be generated by local studies in the developing world, where over 90 percent of aquaculture is practiced and where most wild and captive genetic resources for aquaculture are located. This is part of the global high importance of local studies as contributions to global inferences with respect to fish biodiversity (Palomares et al., 2003). The International Symposia on Genetics in Aquaculture, begun in 1983, contain a wealth of information on aquaculture genetic research and the most important farmed fish species and commodity groups have their associated substantial and ever-increasing bodies of literature on basic research, production, trade etc., including information on breeding programmes and related genetic research results. Good examples are the International Symposia on Tilapia in Aquaculture (ISTAs) (e.g. Fishelson and Yaron, 1983; Bolivar et al., 2004). However, information on FiGR per se in such sources is usually limited and much more is scattered among scientific journals, project reports and other grey literature.

Some of the major contributions to FiGR literature have therefore come from workshops and review papers initiated specifically to collect that scattered information (e.g. Pullin, 1988; Agnèse, 1998; Reddy, 1999; Penman et al., 2005). These mechanisms are useful for compiling information about on-farm, captive FiGR and wild, free-living FiGR. They help to bridge the gap that often exists between mainstream aquaculture literature and mainstream nature conservation literature. For species and commodity groups that are relatively new to aquaculture – often because of very recent advances in technology that allow captive breeding and mass production of seed – information on genetic resources and development of breeding programmes tends to be generated and disseminated more slowly than that for seed production and growout. The current status of sea cucumber fisheries, farming and CBF affords an example (Lovatelli et al., 2004).

FishBase (www.fishbase.org) is the world’s largest biological database on exploited fish, though it covers only finfish. FishBase is constituted and governed as an international consortium of museums, universities and other organizations, including
FAO. Beyond its ongoing contributions to standardization of finfish nomenclature, FishBase contains only limited genetic data of relevance for aquaculture but is still probably the world’s largest compendium of such data in the fields that it has been able to cover so far, including: detailed karyological data for about 200 farmed species; limited electrophoretic population genetics data for about 90 farmed species; and limited quantitative genetics records for 9 farmed species. FishBase also provides online linkages to many other sources of relevant information about aquatic biodiversity, including those emerging as the most important global systems, including the Global Biodiversity Information Facility (GBIF; www.gbif.org) and Ocean Biogeographical Information System (OBIS; www.iobis.org).

FishBase and FAO have provided some information packages on farmed aquatic species, through efforts called respectively “Aquaculture Profiles” and the “Cultured Species Information Programme”. The effort by FAO is ongoing, whereas that by FishBase, begun in the 1990s, has remained stalled for almost 10 years. Table 4 summarizes the results of both, with respect to their choice of species and their coverage of genetic resources, by actual content and/or by pointers to other sources of information. Only 7 of these 32 information packages contain any information on genetic resources per se and only 14 have some links of a limited nature to other sources of genetic resources information.

A new database, “SeaLifeBase”, was initiated in December 2005 to develop for important exploited species of aquatic invertebrates (including farmed crustaceans and molluscs) similar coverage to that provided for finfish by FishBase. SeaLifeBase is being executed from the Fisheries Centre, University of British Columbia, hosted by the FishBase team at the WorldFish Center’s facility in Los Baños, Laguna, Philippines and supported by the Oak Foundation. Under its auspices, representatives of global and regional biological databases, including some that cover farmed or potentially farmable aquatic organisms (e.g. for seaweeds, Algaebase; for some crustaceans, www.crustacea.net; for finfish, FishBase; and for some molluscs, www.data.acnatsci.org/obis/) met from 25 to 27 May 2006 at an Aquaspecies Workshop in Los Baños, Laguna, Philippines, to explore greater collaboration, linkages and interoperability, including establishment of a so-called “SeaLife” portal to provide access to all. It will be important for FAO and others providing or seeking information on genetic resources for aquaculture to monitor all such developments in this dynamic field of work.

The world’s major aquaculture organizations and networks are also useful providers of information of genetic resources for aquaculture, but largely in a current awareness mode and not as genetic resources databases. For example, the Network of Aquaculture Centres in Asia-Pacific (NACA; www.enaca.org) provides a good current awareness facility under the heading “Genetics and Biodiversity”. Similarly “oneFish” (www.onefish.org), a web-based information system developed by the Support Unit for International Fisheries and Aquatic Research (SIFAR; www.sifar.org) in partnership with FAO, provides through its aquaculture and aquaculture resources pages a section entitled “seeds and genetic resources”, linking users to important publications and information about ongoing research and donor programmes. The International Network on Genetics in Aquaculture (INGA; www.worldfishcenter.org/inga) is a useful source of information on the application of genetics in aquaculture and on exchanges of germplasm, especially for some farmed carps and tilapias.

There are many other databases and information systems that provide information on aquatic biodiversity, including those accessible via the World Conservation Union (IUCN; www.iucn.org) and the United Nations Environment Programme/World Conservation Monitoring Centre (www.unep-wcmc.org), but none yet addresses adequately the needs of those seeking substantially aggregated and up to date information on genetic resources for aquaculture. In particular, information about
Table 4
Farmed aquatic species for which information packages are currently available through A. FishBase Aquaculture Profiles and B. the FAO Cultured Species Information Programme, with indications of current importance in aquaculture and whether these sources contain and provide links to genetic resources (GR) information. "NR" = no reliable production statistics available.

<table>
<thead>
<tr>
<th>A. FishBase Aquaculture Profiles</th>
<th>Production year(s)</th>
<th>(t)/countries</th>
<th>+ or - GR info</th>
<th>+ or - GR links</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tilapias:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tilapia rendalli</td>
<td>843 (1995)</td>
<td>8</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Sarotherodon melanotheron</td>
<td>NR</td>
<td>5</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Oreochromis shiranus</td>
<td>ca.10 to 20</td>
<td>1</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Carps:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Labeo rohita</td>
<td>NR</td>
<td>13</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Cirrhinus cirrhous</td>
<td>NR</td>
<td>11</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Catla catla</td>
<td>NR</td>
<td>11</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Others:</td>
<td>Chanos chanos</td>
<td>371,075 (1995)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. The FAO Cultured Species Information Programme</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Seaweeds:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Laminaria japonica</td>
<td>4 917 788 (1999)</td>
<td>4</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Molluscs:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mytilus edulis</td>
<td>NR</td>
<td>12</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Mytilus galloprovincialis</td>
<td>NR</td>
<td>18</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Perna canaliculata</td>
<td>&gt; 70 000 (2002)</td>
<td>1</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ostrea edulis</td>
<td>6-7 000 (2002)</td>
<td>12</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Mercenaria mercenaria</td>
<td>&gt; 40 000 (2002)</td>
<td>3</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Crassostrea virginica</td>
<td>NR</td>
<td>3</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ruditapes philippinarum</td>
<td>236 000 (2002)</td>
<td>9</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Crustaceans:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Macrobrachium rosenbergii</td>
<td>&gt; 200 000 (2002)</td>
<td>12</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Penaeus monodon</td>
<td>676 000 (2001)</td>
<td>21</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Carps: Hypophthalmichthys molitrix</td>
<td>4 100 000 (2003)</td>
<td>6</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Aristichthys nobilis</td>
<td>1 722 832 (2002)</td>
<td>24</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ctenopharyngodon idella</td>
<td>3 572 825 (2002)</td>
<td>&gt;44</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Cyprinus carpio</td>
<td>4 639 460 (2002)</td>
<td>Many</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Carassius auratus</td>
<td>1 702 778 (2002)</td>
<td>14</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Other finfish:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acipenser baerii</td>
<td>NR</td>
<td>12</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Anguilla anguilla</td>
<td>ca. 9 000 (2002)</td>
<td>20</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Argyrosomus regius</td>
<td>231</td>
<td>2</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Dicentrarchus labrax</td>
<td>57 000 (2002)</td>
<td>16</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Ictalurus punctatus</td>
<td>270 000 (1996 for USA only)</td>
<td>6</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Oncorhynchus mykiss</td>
<td>ca. 500 000 (2002)</td>
<td>64</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Psetta maxima</td>
<td>ca. 5 000 (2002)</td>
<td>10</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Salmo salar</td>
<td>&gt;1 000 000 (2002)</td>
<td>17</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Sparus aurata</td>
<td>ca. 90 000 (2002)</td>
<td>18</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Amphibians: Rana catesbiana</td>
<td>NR</td>
<td>13</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Fish breeding programmes, the status and performance of fish strains, hybrids and other genetically altered forms, and fish gene banks is scattered and of highly variable quality; ranging from unverified claims by private breeders to thoroughly documented national collections (e.g. for common carp in Hungary; Bakos and Gorda, 2001). The CABI Compendium on Aquaculture (www.cabi.org/compendium/ac/index.asp) contains useful summaries on major topics concerning genetics in aquaculture and for some species (e.g. Crassostrea gigas and Cyprinus carpio) its coverage extends to and well referenced summaries that include genetic resources information. However, this coverage does not yet extend to all important farmed fish species; e.g. Nile tilapia. As with the abovementioned attempts by FAO and FishBase to provide aquaculture species profiles, all such efforts face the problem that different authors choose to give
different emphases to aquaculture genetics in general and to genetic resources for aquaculture in particular. Moreover, such summaries require frequent updating in order to provide current information in the fast moving field of aquaculture genetics.

For farmed fish, there is not yet any authoritative publication, comparable to the World Watch List for Domestic Animal Diversity (Scherf 1995) from which reliable world totals of breeds and information on their status can be obtained; neither are there any databases for FiGR comparable to those available online for FAnGR: the FAO — maintained Domestic Animal Diversity Information System (DAD-IS; http://dad.fao.org/home.htm) and the International Livestock Research Institute - maintained Domestic Animal Genetic Resources Information System (DAGRIS; http://dagris.ilri.cgiar.org/dagris/). In vitro technologies, especially cryopreservation of fish sperm, are likely to become more widely used for FiGR conservation, as long-term and working gene banks. This will increase the need for online databases through which information about these FiGR can be accessed (e.g. see Kincaid, 2000). The Systemwide Information Network on Genetic Resources (SINGER; http://singer.grinfo.net/) of the Consultative Group on International Agricultural Research currently performs this role for PGR, but not for other genetic resources.

Because of these large gaps for information on FiGR, and because remedying them would assist progressive coverage of genetic resources for aquaculture by the FAO and others, proposals were made, meetings held and initial studies done towards a new information network – initially given the working title “Aquatic Animal Diversity Information and Communication System (AADIS)” and later called a “Fisheries Information Network for Genetic Resources (FINGER)” (FAO, 1999; Pettman, 2002; Pullin et al., 2000, 2002). This initiative has not been taken further, and a fresh approach would now seem more desirable in view of the increased capabilities and interoperability of existing global and regional databases and information systems.

The main growth area in information on genetic resources for aquaculture is that of molecular genetics. More and more information about genetic resources will be in the realm of bioinformatics and not at the species level. This already applies to some farmed populations (e.g. Siraj et al., 1998) and to the huge literature on the genetics of wild populations, especially for salmonids where it is greatly assisting conservation efforts as well as leading to better standardization of criteria and indicators (e.g. Waples et al., 2001; Graziano et al., 2005; Verspoor et al., 2005; Utter, 2004). An “SeaLifeBasees Initiative; FISH-BOL (http://barcoding.si.edu/AllFish.htm) is contributing to the global efforts towards ‘barcoding life’ for all animal species, based on DNA comparisons for cytochrome c oxidase 1 (www.barcodinglife.com). The main challenge with respect to all bioinformatics is to keep as much information as possible in the public domain and accessible to those in the developing world who need it most. This requires further closing of the digital divide between rich and poor nations.

7. THREATS

7.1 To free-living populations

The world’s free-living populations of aquatic species are among its most threatened. Freshwater and diadromous finfish are the world’s most threatened species of high importance to humans. Froese and Torres not cited (1999) found that fishes that depend upon freshwater at any stage within their life cycles are 10 times more likely to be threatened than marine or brackishwater species. In 1998, the increasing global threats to finfish, including many species of importance in aquaculture, were the rationale for a major conference convened by the World Fisheries Trust (Harvey et al., 1998). Cowx (2002) ranked recent threats to freshwater fish as follows: alien species introductions; dams and weirs; water quality problems; habitat degradation; overfishing; flow regulation; overabstraction; tourism; mineral extraction; land use change; climate
change; predators; poor legislation; and “naïve economic criteria”. Freshwater finfish account for at least 65 percent of the world’s production of farmed finfish and some of the world’s free-living populations of freshwater finfish also comprise its most threatened FiGR for aquaculture, not only in terms of the wild relatives of currently farmed species but also for other species that are potential new candidates for aquaculture or contributors to breeding programmes and related research. Tilapia in Africa (e.g. Piers, 2002) and Chinese carps in the PRC (e.g. Wu, 2003) are examples of major groups of threatened genetic resources for farmed freshwater fish.

The world water crisis poses some constraints for expansion of inland aquaculture and for management of some of its free-living genetic resources, but also offers some opportunities for multipurpose use of scarce water resources, adding value to them and benefits from them. Aquaculture can often be an occupier of water rather than a consumer of water. These potentials remain largely unexplored. Most reviews of the world water crisis emphasize domestic water supply and restrict consideration of the importance of water for food production to its use for growing crops. Where fish are mentioned at all in water resources policymaking, this is usually in respect of allowing for some water to remain available for maintaining aquatic ecosystems and biodiversity, rather than recognizing the huge current contributions and scope for future growth of freshwater food fish aquaculture. Where water scarcity is great, however, threats to free-living FiGR are often unavoidable, as illustrated by the following communication to a tilapia genetics list server (L. Kaufman; February 25, 2006; tilapia@lists.unh.edu):

“…the current drought could be threatening the critical refugium populations of Oreochromis esculentus and Oreochromis variabilis in the Lake Kyoga Basin north of Lake Victoria…………Many are assuming that O. esculentus is secure because of the introduced population in Nyumba ya Mungo reservoir, but there is substantial genetic differentiation among the various relict and introduced populations that should not be squandered”.

7.2 To captive populations
Crop and livestock farmers typically discontinue their use of many lower yielding, traditional and minor varieties breeds, for obvious commercial reasons. Their future availability for use in future research and breeding programs is therefore often threatened. For example, 22.5 percent to 32 percent of the world’s livestock breeds are thought to be at risk of extinction (Drucker et al., 2001; FAO data). The same will apply increasingly in aquaculture, as genetic improvement proceeds. Fish seed producers and farmers will choose to keep mainly or exclusively the latest available strains, hybrids etc. The present extent of this has not been documented, but recent indications of wide adoption of GIFT- and GIFT-derived Nile tilapia strains (ADB, 2005b) suggest that it can be rapid.

7.3 Biosafety and biosecurity
For the near future, selective breeding will probably continue to be the main route to genetic improvement in aquaculture. However, increasing use of biotechnology in aquaculture will increase and will involve both use of and impacts upon FiGR and other biodiversity. It must therefore be approached with high precaution and thorough appraisal prior to commercial use. This is biosafety, in the broad sense and it applies to all farmed aquatic organisms, not only to transgenes. As was agreed at a landmark international meeting (ICLARM-FAO, 1999) the characteristics of any genetically altered farmed aquatic organism and its possible impacts on any recipient environments and biota, on-farm and off-farm, are the important biosafety considerations, not the technique(s) by which it was produced.
Despite the high and increasing importance of aquaculture, no farmed aquatic organism has yet been accorded sufficient priority for genome sequencing. There is a strong case for the Nile tilapia genome to be the first farmed fish genome to be sequenced, as this species has global importance in aquaculture and also serves as a model perciform fish (www.hcgs.uhn.edu/cichlid). Development of transgenic fish is well underway (e.g. see http://www.pewagbiotech.ord/research/fish/). Other genetically altered fish, developed from alien and indigenous species, are widely farmed already; for example, highly selected strains, hybrids, artificial polyploids, and monosex populations.

Pullin et al. (in press) found that the proportions of world aquaculture production derived from alien species decreased from about 25 percent in the 1950s to about 15 percent in the 1990s, but pointed out that these data are highly influenced by the huge quantities of indigenous carps farmed in the PRC. On a per country basis, they found that contributions of alien species increased from about 40 percent in the 1950s to 45 percent in the 1990s and that the numbers of alien species used in aquaculture totaled about 40 and were increasing. De Silva et al., (2005), in assessing the roles of alien species in Asian freshwater aquaculture to 2002, found that they accounted for over 40 percent of total production based upon data that excluded indigenous carps farmed in the PRC. With PRC data included, their contribution dropped to almost 12 percent. Casal (2006), from FAO and FishBase data for 2000, found that alien species accounted for only 5 percent of the PRC’s farmed freshwater fish production of 13 269 693 tonnes, but accounted for 72 percent of the 338 861 tonnes of farmed fish produced in Indonesia and 87 percent of the 94 844 tonnes produced in Brazil. It is certain that the use in aquaculture of alien species and of genetically altered forms of both alien and indigenous species will increase. The rapid growth of the farming of Nile tilapia and tilapia hybrids in Asia and Latin America, all developed through original introductions from Africa, and the use of alien Asian species within Asia itself are clear evidence. Consequently, there will be increased movements of farmed aquatic organisms, for production, processing and marketing, as well as for research. This will increase the need for assurance of biosafety, with more effective quarantine and other biosecurity measures. For example, their absence or ineffectiveness and the consequent spread of viral diseases have cost shrimp farming dearly – e.g. white spot syndrome virus, one of four viruses responsible for losses of the order of billions of dollars, cost shrimp farming in Asia (US$4-6 billion) from 1992 to 2001 – and made biosecurity in shrimp farming a growth industry (Lightner, 2005). Specific pathogen-free populations of the Pacific white shrimp (L. vannamei) are becoming genetic resources of importance for shrimp farming.

When aquatic plants and animals escape, or are released for CBF, from research or production facilities, they can have serious adverse impacts (interbreeding, competition for food and for spawning sites, spreading disease etc.) on other aquatic organisms, wild and farmed, and can cause permanent changes to the recipient ecosystems. This applies not only to farmed alien aquatic species but also to farmed genetically altered forms of indigenous species. International introductions, transfers within States, and releases for CBF can bring about permanent changes in the status and integrity of other biodiversity and indeed of other genetic resources for aquaculture. The inevitability of increased use of alien species and of a wide range of genetically altered forms in aquaculture therefore increases the urgency for action to undertake long-term conservation measures for important free-living populations of the wild relatives of farmed aquatic organisms and other species of current or potential importance for aquaculture and related research (see 8.c. below).

Recent meetings and declarations indicate that international and national awareness of the need for biosafety and biosecurity is increasing (e.g. NACA/FAO, 2000; WorldFish Center, 2002, 2003; Gupta et al., 2004). However, moving from such declarations to effective countermeasures against current threats and to ensuring
more responsible future behaviour among actors involved in aquaculture research and development and the entire aquarium trade is not easy, in developed and developing countries alike. Economic growth is the main basis of development and is almost always antagonistic to fish conservation, as shown recently for the USA in a series of papers and a debate led and published by the American Fisheries Society (Czech et al., 2005). Economic growth almost invariably results in widespread losses and degradations of aquatic habitats and reduced aquatic biodiversity.

8. MANAGEMENT

8.1 Concepts and definitions
Management of aquatic genetic resources is full integration of their use and conservation (Pullin, 2000). Conservation of FiGR of actual or future potential use is itself a form of use. Genetic resources can be conserved by one or more of the following options: in situ/in vivo, as captive or free-living populations; ex situ/in vitro, as gametes, embryos, other tissues and DNA; and ex situ/in vivo, as captive populations in research establishments, aquaria, etc.

The Convention on Biological Diversity (CBD 1994) definitions for genetic resources and related terms are followed here, as they are in the mainstream PGR and FAAnGR literature. In most FiGR literature, however, use of the terms in situ and ex situ to describe FiGR is not yet consistent with CBD definitions. According to the CBD (1994), in situ conditions are those “... where genetic resources exist within ecosystems and natural habitats, and, in the case of domesticated or cultivated species, in the surroundings where they have developed their distinctive properties”; and ex situ conservation is “conservation of components of biological diversity outside their natural habitats”.

This means that the genetic resources of farmed aquatic organisms that have distinctive properties, and that are held in vivo (i.e. as live populations) in their typical farm environments should properly be called in situ, as should all wild and feral genetic resources for aquaculture in their typical habitats. The term ex situ should be used only for FiGR and aquatic PGR held in vitro (e.g. collections of cryopreserved fish spermatozoa, embryos and other tissues) and for FiGR and aquatic PGR held in vivo in artificial, off-farm environments (e.g. botanical gardens, aquaria, research establishments and zoos). However, for captive fish populations, the distinction between typical farm environments and these atypical off-farm environments cannot yet be applied as strictly as it can for crop varieties and livestock breeds.

The CBD does not define or elaborate on “distinctive properties”. However, for broad categorization of wild and captive genetic resources for aquaculture, it can be assumed that all captive-bred populations of farmed aquatic species have undergone some genetic alteration so as to differ genetically from free-living populations of the same species. The degrees of genetic alteration vary greatly according to with the different histories of farmed aquatic populations with respect to artificial selection, interstrain, interspecifics and intergeneric hybridization, as well as genetic manipulations, including control of sex determination, artificial polyploidy, androgenesis, gynogenesis and transgenesis. Irrespective of all of these purposeful interventions, all captive populations undergo natural selection to hatchery and farm environments.

8.2 In situ/in vivo; captive populations on-farm
In situ/in vivo conservation of FiGR on farms is accomplished mostly by seed producers, as broodstock populations. However, there are narrow limits to the diversity of FiGR that can be conserved and used by commercial seed producers and farmers. They must use the bulk of their facilities for holding and selling fish of highly proven
viability and profitability, unless compensated specifically to keep other species and strains for conservation purposes. The same applies to FAnGR, for the conservation of traditional and rare breeds of livestock on-farm.

The main requirements for most conservation of FiGR as broodstock on farms, and indeed as ex situ/in vivo populations in other facilities (see 7.e. below), are acquisition of founder stocks with high genetic variance and thereafter maintenance of adequate breeding numbers, so as to avoid inbreeding. Broodstock are often not managed well, especially in developing countries. The temptation is to keep only small effective breeding numbers of highly fecund species, such as farmed carps, and to practice ad hoc replacement of far less fecund tilapia broodstock from whatever sources are available. Broodstock replacement is expensive. For example, tilapia broodstock used for seed production should normally be replaced within two years of the start of their productive life.

Excellent guides are available for broodstock management and for the selective breeding that it facilitates (e.g. Tave, 1986, 1989; WorldFish Center, 2004a; Gjedrem, 2005). Where farmed fish breeding programmes are well developed, government ministries and research organizations, fish producers associations, certified private sector breeders, and farmers can all work in concert to conserve valuable breeds and to maintain seed quality; for example, in Hungary, 13 breeding farms of the Carp Breeding Section of the Hungarian Fish Producers Association keep 24 certified common carp strains (Váradi et al., 2002).

8.3 In situ/in vivo; free-living populations

Free-living, wild and feral, populations of farmed and potentially farmable aquatic species, in inland, coastal and marine waters and wetlands, comprise genetic resources of immense importance for aquaculture. For example, Pullin et al., (2001), from FishBase data, found among the fish fauna of Africa 2 608 unique freshwater species and 842 unique marine species, with over 100 fish species being used in aquaculture and over 1 000 in the aquarium trade. Information about the genetic diversity of some of their populations is increasing together with efforts for their conservation (e.g. Agnèse, 1994, 1998; Ryman et al., 1995; Lévêque, 1997; Miller and Craig, 2001; Collares-Pereira et al., 2002; Abban et al., 2004), but the genetic diversity of many is still very imperfectly known. For example, local populations of marine organisms, particularly invertebrates, can exhibit high levels of cryptic speciation (Thorpe et al., 2000).

Conservation of important free-living FiGR is essentially nature conservation. It depends upon the maintenance of their habitats and prevention of human influences that could cause genetic change, including isolation from aquaculture development, alien species and genetically altered farmed aquatic organisms. Aquatic protected areas can provide this to some extent, though conservation of FiGR for aquaculture is still seldom mentioned as a major reason for their establishment, relative to other reasons given: e.g. increased recruitment of neighbouring capture fisheries (e.g. not cited Shiple, 2004). Moreover, far greater emphasis has been given so far to marine protected areas than to freshwater protected areas for the more important and threatened FiGR for freshwater aquaculture. As Rice (2005) has pointed out, managing fish habitats for conservation purposes must keep pace with the rapid scientific developments and new thinking about ecosystem management. Habitat science per se has so far lagged behind ecosystem science.

Pullin (1990) recommended increased emphasis on conservation of fish genetic diversity among the goals of nature reserves and safari parks but, as with protected areas in general, this would not often guarantee the high degree of isolation needed to prevent disruption and genetic change. Important PGR are conserved in relatively small areas of habitats that are kept pristine or near-pristine as sacred groves etc.
(e.g. Okafor and Ladipo, 1992) and the extents to which FiGR are also conserved at such locations should be documented. For a more widely applicable and essentially new strategy, Pullin (2006b) suggested co-financing the establishment and upkeep of FiGR reserves, permanently isolated from all contact with aquaculture and other disturbances, together with the responsible development of other areas of aquatic ecosystems, including aquaculture development.

8.4 Ex situ/in vitro; cryopreserved sperm, embryos and tissue/DNA banking

*In vitro* cryopreservation of fish sperm has been accomplished for many species (Tiersch and Mazik, 2000) and is probably achievable for all farmed fish, though frozen sperm viability varies greatly with species. Cryopreservation of the early embryos of bivalve molluscs and sea urchins is also technically possible. However, the large size and fragility of most finfish eggs and embryos have so far defeated all attempts at their cryopreservation. Despite widespread successes with cryopreservation of farmed fish sperm at aquaculture research institutes and fish breeding centres around the world (for examples, see papers in Harvey *et al.*, 1998), this technology remains little used by fish breeders and seed producers, especially in developing countries. It is the obvious future mainstay for long-term, *in vitro* gene banking of FiGR for aquaculture, including farmed and potentially farmable fish, their wild relatives, and all other to in situ/in useful and potentially useful fish genetic material. Savolainen *et al.* (2006) have reviewed prospects and practices for banking DNA and tissues. This has been conceived mainly for plants, but could be explored for farmed aquatic animals.

*Ex situ/in vitro* conservation of FiGR is best viewed as complementary to their *in situ/vivo* conservation, as has been the strategy for most of the world’s PGR. The World Fisheries Trust (www.worldfish.org) has long pioneered complementary conservation of FiGR as free-living populations and as cryopreserved fish sperm, and undertakes extensive training for this approach in developing countries.

8.5 Ex situ/in vivo; captive populations in aquaria and research establishments

Public and private aquaria have great scope for conserving FiGR, but this has not yet been realized to the extent of the role played by zoos in conservation of FAnGR. Wild relatives and rare breeds of livestock in zoos are often managed not only as public exhibits but also as *in vivo* gene banks. The population genetics of farmed fish held in aquarium collections have been little studied. Public and private aquaculture research establishments already play large roles in conservation of farmed fish, as captive populations. The problem here is that maintaining and replacing in vivo fish populations is expensive in terms of facilities, staffing and feeds, fish health care etc. The fish research collections of many universities that undertake aquaculture research and teaching are indeed *in vivo* gene banks, provided that their existence does not end along with the short-term projects for which many accessions are acquired.

The Research Institute for Fisheries, Aquaculture and Irrigation (HAKI) leads Hungary’s National Carp Breeding Programme (CBP), in collaboration with the Common Carp Breeding Section of the Hungarian Fish Producers Association, using standard methodology (OMMI). HAKI keeps an *in vivo* gene bank of over 30 strains of farmed and wild common carp (e.g. Bakos and Gorda, 2001; Bakos *et al.*, 2002) Since 2002, however, the government ceased to provide support for HAKI’s *in vivo* carp gene banking, which HAKI must now fund from its own budgets. Some 25 private farmers maintain populations of their own strains under the CBP. Farmers receive subsidies if they produce OMMI-approved common carp strains (L. Váradi, J. Bakos and Z. Jeney; personal communications).

A further constraint in many developing countries is that tradition or economic necessity requires some government research institutions to produce large quantities
of seed for distribution to farmers. This function can severely limit the availability of facilities for \textit{in vivo} gene banking.

9. SHARING BENEFITS AND COSTS: OWNERSHIP AND USE

9.1 Free-living populations
The CBD gives its Parties national sovereignty over their biodiversity, including FiGR for aquaculture. The CBD also provides for recognition of new countries of origin for populations of farmed aquatic organisms that have acquired distinctive properties outside their native ranges; for example, the distinctive farmed strains of common carp developed in Indonesia. The CBD, together with other international conventions that concern aquatic ecosystems (notably the Ramsar Convention on Wetlands, 1971 and the United Nations Convention on Law of the Sea, 1982) also imposes national obligations on Parties to conserve their living aquatic resources.

Poor countries cannot easily take on the burden of conserving their extensive free-living FiGR for use in world aquaculture without external financial and technical support. Many of the world's important free-living FiGR for aquaculture are owned, and often used, by poor indigenous peoples and local communities, who cannot afford to be their long-term stewards for use by the rest of the world unless adequately compensated. The CBD's Article 8j provides for this in common with other international provisions on human rights (e.g. Posey, 1999). Greer and Harvey (2004) have reviewed some of the limited progress made in implementation of these provisions. There have not yet been any well documented examples of the stewards of free-living FiGR for aquaculture and other users of those FiGR for commercial purposes sharing the costs of conservation and the benefits of use.

9.2 Public and private research
Since the 1980s, developed countries seem to have shifted their public-sector research priorities away from increasing the production of food staples (that, coincidentally, provided useful spillovers to developing countries), putting more emphasis on research on environmental, food safety and various other non-food production aspects of agriculture. This trend may require developing countries to invest more in food production research, becoming more self-reliant (Pardey \textit{et al.}, 2006). At the same time, private research and development of biotechnology for staple food commodities has increased, with a growth in intellectual property rights (IPR) and growing concerns as to how these trends will affect developing countries (Wekundah, 2005; Wright and Pardey, 2006a, 2006b).

Private sector research in biotechnology for aquaculture has also increased, especially in developed countries, and the developing countries where most of the world's fish are farmed will need to increase their own public and private research capacities in this area if they are not to be left behind. However, private ownership of FiGR for aquaculture, through assumption of intellectual property rights (IPR) or other restrictions on use, is still rare. There are no well documented examples of substantial financial returns to researchers who have developed and assumed ownership of specific FiGR for aquaculture and related biotechnology. Ownership rights and restrictions on use of FiGR are usually very difficult to enforce. Farmed fish from different breeding programmes and genetic manipulations often look alike and therefore the provenance of a given farmed fish population \textit{in situ} or in a market place is difficult to determine without costly forensic examinations.

For example, GIFT and GIFT-derived and other improved strains of Nile tilapia all look very similar. Without recourse to laboratory tests, a casual observer of their farmed populations and harvests could say only that they must be genetically improved
rather than unimproved fish. Simpler and cheaper genetic marking of superficially similar farmed fish strains, hybrids and other genetically altered forms will likely become available to help their developers to differentiate between legitimate use by those who have signed restrictive use agreements and others who are enjoying pirate use. However, acquiring and enforcing IPR on FiGR for aquaculture as strains, hybrids and other genetically altered forms will remain difficult. Their complexities are increased by the prevalence of public-private partnerships in fish breeding, seed supply and farming. It is common in some developed and in most developing countries for government research establishments, breeding centres and hatcheries to supply genetic material to the private sector and also to act as substantial producers of fish seed, even though this latter function could take significant market share away from private seed producers. This issue has emerged in the public-private relationships associated with tilapia breeding and seed supply in the Philippines (WorldFish Center, 2004b).

It is worth noting, at this early stage of domestication for most farmed fish, that the main traditional and commercial breeds of livestock and pet animals (e.g. the Holstein cow and the Labrador dog) are not privately or even nationally owned. Rather, there is private ownership of and restricted, purchasable access to the progeny of multiple pedigreed strains and to individual sires and dams. Hamilton (1999) found no instances of attempts to claim even national or regional sovereignty over or controlling interests in any livestock breed. Pedigreed fish populations in a single hatchery or farm, and pedigreed fish sires and dams are still very little developed compared to their prevalence in livestock and pet animal breeding, but their development would probably afford a better basis for the acquisition of private rights to and returns from FiGR than attempts to seek patents or other officially recognized IPR on farmed fish strains and other genetically altered forms.

The main requirement for equitable use of FiGR is better organization and oversight of germplasm acquisition and transfers, through Germplasm Acquisition Agreements and Material Transfer Agreements similar to those developed for PGR. Public, private and public-private transfers of FiGR for aquaculture are increasing. Responsible protocols and practices for these are not yet well developed or enforced. The INGA has contributed to improving this situation.

10. STRATEGIC DIRECTIONS

10.1 Increased investment
The growth of aquaculture has outpaced that of all other food production sectors and its high importance and scope for further growth, especially for the benefit of poor consumers and farmers, are clear. Past and present investment in the management of genetic resources for aquaculture fails to reflect this. If this situation continues, it will jeopardize achievement of the potential of aquaculture. Many genetic resources for aquaculture are seriously threatened. Countermeasures require increased investment in their management, to match their economic and social importance.

Effective management of genetic resources for aquaculture almost always has higher costs than are normally encountered with PGR and FAnGR. Setting aside areas of natural ecosystems as off-limits to all forms of disturbance has operational and opportunity costs. Establishing and maintaining \textit{ex situ}, \textit{in vivo} and/or \textit{in vitro} fish gene banks is very expensive compared the costs involved in plant gene banks, and gene banks for FiGR cannot be centralized to the same extents as those for PGR. National, regional and international networks and partnerships, including public-private partnerships, can help in the sharing of costs for and benefits from management of FiGR for aquaculture. For example, in Central and Eastern Europe, the Network of Aquaculture Centres (NACEE; http://agrowebcee.net/subnetwork/nacee/) links 31 institutes from 13 countries, all having strong interests in carp genetic resources (Bakos et al., 2002).
10.2 Management as agrobiodiversity
The whole of agriculture and fisheries and their supportive ecosystems function as a global trophic web. However, aquaculture is farming and has much more in common with agriculture than with capture fisheries. In particular, on-farm *in situ* and all *ex situ* genetic resources for aquaculture merit recognition as part of agrobiodiversity and management, along with PGR and FAnGR, through common policies, institutions and mechanisms.

10.3 Improved information systems
Thorough documentation and accessible information on all categories of genetic resources for aquaculture is an urgent requirement. This means gathering, processing and linking information on free-living genetic resources for aquaculture with that for breeding programmes and related research, with the types of seed supplied to farmers, and with production and value statistics for farmed aquatic species, strains and other genetically altered forms. This can be approached progressively. The genetic resources of the more important farmed aquatic plants could be covered under existing arrangements between the International Plant Genetic Resources Institute (IPGRI) and FAO. It would also be relatively easy to prioritize coverage of the most important genetic resources for farmed food fish. The genetic resources for farmed ornamental aquatic species are a lower priority and will continue to be documented to some extents by the aquarium trade and by databases such as FishBase.

10.4 Conservation in changing ecosystems
The future availability and integrity of free-living and captive genetic resources for aquaculture depends upon the status of their environments; i.e., natural aquatic ecosystems and agroecosystems. Brown *et al.*, (1997) made this point thus, with reference to pressures such as fragmentation, and pollution: “…..the goal of conserving appropriate genetic diversity is best achieved not by focusing on maintenance of the genes and genotypes that currently exist within a species, but by trying to prevent drastic alteration in the pace and direction of these evolutionary processes.”

This amounts to a call for ecosystem-based management at the genetic level, on-farm as well as for natural ecosystems. The increasing needs to confront climate change and climatic uncertainties are also highly relevant here. However, much of the literature on ecosystem-based management for fisheries emphasizes the species level, higher taxa and their functions, and pays little attention to genetic resources. An ecosystems perspective that includes the genetic level will show that some losses of genetic resources for aquaculture are inevitable as development proceeds. It is important to recognize this and, by monitoring and understanding the processes involved, to improve prospects for keeping important genetic diversity. What actually can be kept and what will be lost are parts of a bigger picture than genetic resources inventories alone can suggest, and the costs of *in situ/in vivo* conservation and complementary *ex situ/in vitro* conservation are always serious constraints. The conservation of free-living populations and traditional breeds of farmed species is like a battlefield where, distasteful though it is, triage is sometimes inevitable. Complementary *ex situ*, *in vitro* and *in vivo*, conservation is vital for important genetic resources that are seriously threatened *in situ*.

10.5 Reconciliation of aquaculture and nature conservation
Conservation of *in situ/in vivo*, free-living genetic resources for aquaculture have yet to be adequately recognized as part of the rationale for greater investment in conservation of natural aquatic biodiversity and habitats. Many nature conservationists can conceive alliances between agriculture or forestry and conservation but most perceive aquaculture principally or solely as a threat. As more responsible aquaculture becomes the norm, the CBD, IUCN and the Ramsar Convention, together with many
nature conservation organizations, especially NGOs, at international, national and local levels, will hopefully find partners within the aquaculture sector itself so as to reconcile and, where possible, to twin their respective goals. FAO and the CGIAR can help this process, but are likely to be more involved with conservation of captive and in vitro genetic resources for aquaculture production and related research.

10.6 Progressive linkages with management of FAnGR
Recent meetings and publications (Pullin 2006b; Science Council 2005) have recognized the many lessons to be learned from management of FAnZGR for management of FiGR for aquaculture. For example, there could be much closer linkages with respect ex situ/in vitro conservation of FiGR and FAnGR, especially in terms of shared facilities. The main strategy for FiGR here would probably be decentralization, with establishment of and support to relatively small and affordable national and local gene banks, kept within or as close as possible to production areas. Most responsibilities would probably rest with national public sector research establishments, private sector breeders and seed suppliers. The CGIAR would probably not be involved to any extent comparable with its involvement in gene banks for PGR. The WorldFish Center has so far taken only a minor role in this area to date, for GIFT strains of Nile tilapia and for its own collaborative and in-house research. The International Livestock Research Institute is not involved in gene banking for FAnGR, but has collections of PGR for fodder species.

10.7 Exploration of interactive governance and governability
Management of genetic resources for aquaculture is part of the global management of all natural resources. A new approach to this, called interactive governance, is being developed, using capture fisheries as its main model, with some preliminary explorations for aquaculture (Kooiman et al., 2005; Bavinck et al., 2005; Pullin and Sumaila, 2005). Interactive governance recognizes the diversity, complexity, dynamics and scales that are represented in all natural resources that are "systems to be governed". Genetic resources for aquaculture fit this description very well and are therefore subjects for further explorations of the utility of the interactive governance governance approach for their management and for assessments of their governabilities. Research in this general area is being carried out by an international network (www.fishgovnet.org) with a current emphasis on operationalizing interactive governance in capture fisheries, aquaculture and coastal zones, mainly through developing the concept of and methodologies for determining governability (e.g. Chuenpagdee et al., 2005).

11. ACKNOWLEDGEMENTS
The author thanks the following for helpful responses during the preparation of this review: János Bakos, Christine Casal, Jean Collins, Peter Edwards, Rainer Froese, Gideon Hulata, Zsigmond Jeney, Alan Lowther, Graham Mair, Phil Pardey and László Váradi.

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Fish genomics and analytical genetic technologies, with examples of their potential applications in management of fish genetic resources

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1. SUMMARY
The successful completion of the human genome project marked the start of a genomics revolution, which has the potential to impact aquaculture and fisheries production and has implications for the future management of fish genetic resources (FiGR). Aquaculture has the potential to compensate for dwindling capture fisheries, but biological and production hurdles must be overcome in order to develop cost-effective and sustainable aquaculture. Diseases are major threats to sustainability, and therefore the disease problems should be given high priority. In addition, important production and performance traits (such as growth rate, feed conversion efficiency, body conformation and fillet yield) must be improved in order to make aquaculture more productive and profitable. Genetic enhancement of farmed fish is needed not only to meet the demands of fish production, but also to ensure profitability.

The analytical genetic technologies most relevant to aquaculture and capture fisheries include: DNA markers, genome mapping, microarrays, and sequencing. DNA marker technologies are not only the basis for genetic linkage mapping, but also for the analysis of genetic resources, strain differentiation, species differentiation, parentage identification, and preservation of genetic diversity and conservation of genetic integrity.

The application of genomics in aquaculture is still at the early stages. For many important species of farmed fish, molecular markers have been developed allowing genetic analysis for FiGR conservation and genetic enhancement of farmed fish. Linkage and physical maps have been developed allowing elucidation of genes responsible for important performance and production traits; genome reagents such as expressed sequence tags have been produced providing material basis for the development of microarray technology.

Studies of the genomes of farmed and fished aquatic species have shown both common and unique characteristics that provide both advantages and challenges. In most cases, the genomes of farmed aquatic species are smaller than or comparable to the human genome. Many farmed aquatic species have high fecundity that provides large full-sib and half-sib families, and this greatly facilitates quantitative trait loci (QTL) mapping. However, the large number of farmed aquatic species tends to dilute genomic research efforts.
The genomics revolution and its impacts on aquaculture are expected to contribute to resolving problems such as diseases, environmental impacts, and low profit margins. The major potential applications of genome technologies, primarily in aquaculture but also to some extents in capture fisheries include: marker-assisted selection (MAS) for genetic enhancement; environmental improvements through enhanced productivity as well as the development of novel technologies for environment monitoring, development of effective vaccines and their delivery technologies; monitoring antibiotic resistance; diagnosis for fish diseases and for the safety of aquatic produce; accurate identification of fish stocks for capture fisheries management and for their use as FiGR in aquaculture; conservation of FiGR, including protection of endangered species, in response to fish production strategies and consumer interests; and the development and application of transgenic fish technology including, for example, sterilization technology to address concerns about their possible environmental impacts.

A great challenge for aquaculture and capture fisheries is the long-term conservation of FiGR. Genome technologies provide new tools for genetic analysis. Innovative DNA marker technologies have opened a broad avenue for the analysis of genetic diversity based on genotypes. Some aquaculture operations still use wild fish seed. For these and for future fish breeding programs, conservation of wild FiGR is important.

The applications of genomics in aquaculture and capture fisheries raise ethical, economic, environmental, legal, and social concerns. The most prominent of these at present relate to the development and use of genetically modified organisms. More research is needed not only to resolve issues related to safety of using transgenic fish, but also to produce novel technologies allowing safe use of transgenic technology.

Public education about genomics and its applications is a key issue. The public is relatively naïve and ill-informed about genomics. Conversely, genomics researchers may not understand the practical needs of aquaculture and capture fisheries or of fish consumers. While information dissemination about genomics to the public is very important, better exchanges of information between genome researchers and aquaculture and fisheries professionals are also essential.

Fish genomics and analytical genetic technologies are reviewed here, with some examples of their implications for FiGR management. Genomics is a highly dynamic research field, currently dominated by human genomics but rapid developments in genomics can afford new opportunities for applications in aquaculture and capture fisheries, particularly in the areas of FiGR conservation and genetic enhancement.

2. BACKGROUND
Genomics began to receive substantial attention as a result of the Human Genome Project. The Human Genome Project faced the tasks of decoding the three billion base pairs of the human genome. Genomics always generates large data sets and these demand new ways of data management. Genomics draws data from cytogenetics, molecular genetics, quantitative genetics, and population genetics, and has led to the development of bioinformatics, through which raw genome information links to meaningful biological information. Genomics comprises the study of genome structure, organization, expression, evolution, and functions. Many sub-branches of genomics are emerging, including aquaculture genomics (http://www.genomicglossaries.com).

Genomes and genomics
The term genome refers to the complete genetic material of an organism. This includes the nuclear and mitochondrial genomes for plant and animals, and also chloroplast genomes for plants. Mitochondrial and chloroplast genomes are small and contain only a limited number of genes. The focus of most genome research is on the nuclear genome,
through mitochondrial genomes have been extremely useful for the identification of fish species and populations. Genomics is the science that studies the genome.

The genetic information stored in DNA cannot be used without being transcribed into RNA which then, with very few exceptions, must be translated into proteins in order to have biological functions. The term genomics often is used to cover not only this narrow sense genomics, but also transcriptomics, and in many cases proteomics as well. As Figure 1 shows, the entire DNA content of an organism (the genome) is transcribed into RNA (the entire RNA content of the organism is called the transcriptome), and the RNA is translated into proteins (the proteome). Genomics, transcriptomics, and proteomics are sciences that study the genome, transcriptome, and proteome, respectively. Genomics can be divided into structural genomics, which studies the structures, organization, and evolution of genomes, and functional genomics, which studies expression and functions of the genomes.

Genetic diversity at the genome level
Through the long process of evolution, many mutations and other genetic changes have accumulated. Accumulation of different mutations in reproductive isolated populations and individuals, as a result of their environment, is the fundamental basis of fish genetic diversity. The basic idea behind fish population genetic analysis is to unravel such differences and their inheritance among populations.

Whereas the genome is relatively stable in an organism, the transcriptome is highly dynamic. The types of transcripts and their relative levels of expression are highly regulated by tissue specificity, developmental stage, physiological state, and the environment. For instance, an organism might have 25,000 genes, but not all are expressed in every type of cell. Those genes required for the basic cell structure and functions are probably expressed in all tissues, organs, and cell types; whereas each cell type expresses a subset of the genes specific for those cell types. Many genes are expressed throughout the life history of an organism, but certain genes are expressed only at a specific developmental stage. The environment can insert its effect on gene expression in multiple dimensions. Temperature, pH, water quality, stress, dissolved oxygen, and many other environmental factors can induce or suppress expression of a large number of genes. Environmental pollution can lead to activation and suppression of expression of many genes in both the types of genes being expressed, and the levels of gene expression. Consequently, genome technologies have much to do with the environment, as well as the genome. It is now widely believed that the complexity of the transcriptome is much larger than the genome, because of alternatively processed transcripts. The information stored in the genome is amplified and diversified once at the RNA level, and is further amplified and diversified at the protein level by post-translational glycosylation, acetylation, phosphorylation, and other modifications leading to drastically different biological functions.
3. THE GENOMICS REVOLUTION AND ITS EMERGING TRENDS

Francis Collins, the Director of the National Human Genome Research Institute (NHGRI), with inputs from 600 scientists, described the “three-floor house” for the future of genomics (Box 1).

From this three-floor house plan of genomics, the following trends can be deduced:

- Genomics goes functional
- Genomics goes global
- Genomics will continue to be dominated by human genomes and human health concerns
- Genomics goes environmental
- Genomics moves towards systems biology, metagenomics, and predictive biology
- Advances in genomics will accelerate; some significant discoveries and their impacts will probably be unintentional

Genomics goes functional

The first and the overwhelming trend of genomics is that it is going functional. Although having the human genome sequenced still seems new, genomics research is rushing ahead to the next step, functional genomics. What functional genomics covers depends largely on who is being asked, but many scientists agree that the scope of functional

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**BOX 1**

The three floor house of future genomics

**First floor: genomics to biology**
- Define the structure of genetic variations with tools like the HapMap.
- Decode many additional genomes.
- Reduce the cost of decoding an individual genome from $30 million today to $1,000.
- Identify the functional elements of the human genome.
- Identify all proteins of the cell and their interactions.
- Build a computational model of a human cell and extend it to many types of human cells.

**Second floor: genomics to health**
- Identify genetic and environmental risk factors for common diseases.
- Build “sentinel” systems to detect diseases before they become advanced.
- Get academic researchers to embrace modern drug discovery techniques to create promising compounds.
- Gather and study genotypes from a 500,000-person segment of the U.S. population for 7 to 10 years.
- Figure out modern-day health disparities such as why some groups are afflicted with specific diseases more than others.
- Use genomics outside our borders to combat malaria, tuberculosis, and other diseases.

**Third floor: genomics to society**
- Enhance genetic privacy and protection via legislation.
- Understand genomics with respect to race and ethnicity. Be wary of demagogues who would exploit genomics for political purposes.
- Understand genomics’ impact on human characteristics such as intelligence and sexual orientation.
- Define areas that should not be exploited, such as designing characteristics of future generations.

Genomics ranges from expression profiling, the relationship between genome expression and functions, discovery of gene functions and their interrelationships, understanding networking among genes in relation to carrying out their functions, to proteomics and protein-protein interactions. Potential application areas include clinical diagnostics, agro biotechnology, environmental biotechnology, and pharmacogenomics. Although functional genomics remains young enough that people argue over their definition, few squabble over the value of this field. Advances in areas from gene expression to proteomics promise to push ahead basic research, biotechnology, and medicine. In fact, some experts predict an annual compound growth rate of 28% for the next six years in commercial sectors of functional genomics. As functional genomics moves forward, it will provide many options for applications in aquaculture and capture fisheries.

Genomics goes global
Genomics is going yet more global as many countries have an interest in participation, and no one wants to fall behind. Genome science is so big that no single individual or single laboratory can do it alone. It requires collaborations, team work, and international cooperation. Not only is international cooperation important for genome research because many of the genetic resources are shared by the international communities, collaboration among the private sector is another trend. The complexity concomitant with genomics and proteomics has had two key organizational impacts for large pharmaceutical companies (Arlington and Peakman, 2001). First, it has created a situation wherein the industry no longer has the resources to cover every technology, disease and therapeutic area. The second impact of the genomics revolution is to lower the entry barriers to new competitors who might be much more nimble in finding and validating targets and leads using virtual networks. These changes are per se healthy from the perspective that more opportunities are created for new players, while existing giant players have to ask the hard question about how to maintain a competitive edge in the genomics era with explosive growth with the amount of available information. Companies and national programs need to consider making adequate investments in education and capacity building to provide the human resources needed to take genomics forward.

More consortia are likely to be established to address the big questions that genomics can answer, but such questions are too big and too risky for companies to tackle on an individual competitive basis (Arlington and Peakman, 2001). Aquaculture research communities are already working collaboratively by forming various genome consortia including Salmonids Genome Consortium, Oyster Genome Consortium, and Catfish Genome Consortium etc. (e.g., http://web.uvic.ca/cbr/grasp/). The Animal Genome Project in the United States of America is organized under a National Project of NRSP-8, in which aquaculture genome is a component. Under NRSP-8, each species has a coordinator (http://www.animalgenome.org/).

Genomics will continue to be dominated by human genomes and human health concerns
Human genomics and human health concerns will continue to dominate genomics, even though many other areas, such as agricultural genomics and environmental genomics may be equally important because they affect human health. All of the genomics information and genetic technologies developed to date will be exploited to the maximal extent in human health and pharmaceutical developments. Genome technologies and genomic information allow genetic testing to be performed with a much greater precision for the prediction of predisposition to disease and ailments, carrier status, and prenatal testing. Such capabilities likely will lead to a trend of using genome technologies for pre-symptom predictions of diseases. Genome information,
genome resource, and genetic technologies also assist development of pharmaceutical products, including genes and gene products.

**Genomics goes environmental**

Public concern about environment changes and environmental quality is high and there are driving forces to address this in government, NGOs, and the public at large, with modern technologies available for the benefit of the environment (Gracey and Cossins, 2003; Cossins and Crawford, 2005; Almeida et al. 2005). Such technologies can be grouped into two general categories: those that enhance agricultural production with the same or less input, and those that can provide novel sentinels for environmental monitoring. For instance, broodstocks can be selected for better feed conversion efficiencies using gene-assisted technologies allowing greater yields with less feed, reducing environmental problems from agricultural production including aquaculture; microarrays can provide precise information on environmental pollution and its impact on the organisms involved in the system. Functional genomics is expected to contribute information for defining environmental issues, as well as technology for environmental monitoring and environment-friendly technologies for agriculture, aquaculture, and natural resource utilization.

**Genomics moves towards metagenomics, systems biology, and predictive biology**

Genomics is moving toward systems biology, metagenomics, and predictive biology. Genomics, once wholly described by single-organism sequencing efforts, is poised to fulfill its scientific promise in a number of different ways as sequence information is transformed into biological meaning by evolving technologies, theoretical frameworks and practical goals. Systems biology and metagenomics are two of the most ambitious of these emerging genomic sciences, concerned with 'total' understanding of cellular and ecological systems. Metagenomics is also referred to as environmental genomics or community genomics. It is the culture-independent genomic analysis of microbial communities (Eyers et al., 2004; Galperin, 2004; Riesenfeld et al., 2004; Rodriguez-Valera, 2004; Schloss and Handelsman, 2003). Systems biology aims to reconcile the exponentially growing amount of data about macromolecules, cells, tissues, organisms, populations, and ecosystems into coherent and systemic views of organization (Ge et al., 2003; Kitano et al., 2002). The genomics era has led to a much greater understanding of physiology and pathology at the molecular level and is enabling scientists to begin to unravel cellular processes as the result of the interplay of networks of genes. The publication of the human genome sequence and the use of expression databases and sophisticated bioinformatics software to find and characterize new genes and gene families have identified a huge number of potential and actual targets in a wide range of diseases. Further, the understanding of genome variation and the impact this has on health and disease will significantly improve the development and delivery of new medicines. On top of systems biology, modern genome sciences should generate information concerning expression of genomes as to “when this happens, then that happens”. Predictive biology will provide insights as to whether and when certain conditions, such as disease epidemics, may or may not happen.

**Advances in genomics will accelerate: some significant discoveries and their impacts will probably be unintentional**

With the great expectations from the human genome project and the potentially revolutionary advances of sequencing technology, it is likely that genomics will make new rapid leaps forward. While sequencing a single genome was regarded as utopia 20 years ago, sequencing thousands of human genomes is now possible. Such sequencing
capacities and efficiency, when coupled to the ability to analyse the genomics data and to disseminate them through bioinformatics, suggest that there will be great advances in genomics. However, some discoveries and their impacts will probably be unintentional. Computerized analysis of complex genomics data can bring discoveries that are not related to the main purposes for which they were collected.

4. EXAMPLES OF FISH GENOMICS AND ANALYTICAL GENETIC TECHNOLOGIES
Since the completion of the Human Genome Project, major progress has been made in genome research, including the genomics of some farmed fish. The first Workshop on Aquaculture Species Genome Mapping was held in May 1997 in Dartmouth, Massachusetts, United States of America. Thus it was decided to focus on five species groups in the United States of America: catfish, tilapia, salmon/trout, shrimps, and oysters. In 2003, an Aquaculture Genome Project joined USDA project NRSP-8, as a part of the National Animal Genome Project, with the addition of striped bass (*Morone saxatilis*) as the sixth aquaculture species of focus in the United States of America.

The most potentially useful genomic and analytical genetic technologies for application in aquaculture and capture fisheries are: DNA markers, genome mapping, and microarrays (see Annex 1). DNA marker technologies include various techniques and methods for the analysis of genetic variation at the individual, population, or species levels. They are not only the basis for genetic linkage mapping, but also for the analysis of genetic resources, strain differentiation, species differentiation, parentage identification, and preservation of genetic diversity and conservation of genetic integrity.

**Examples of major aquaculture genome projects**
The US NRSP-8 project was initiated in 1998 and it is now in its second five-year phase (2003-2008). The major objective in the first phase was to develop molecular markers, and construction of genetic linkage, physical, and radiation hybrid maps. The project has three objectives in its current phase: 1) enhance and integrate genetic and physical maps of agriculturally important animals for cross species comparisons and sequence annotation; 2) facilitate integration of genomic, transcriptional, proteomic and metabolomic approaches toward better understanding of biological mechanisms underlying economically important traits; and 3) facilitate and implement bioinformatic tools to extract, analyze, store and disseminate information (http://www.animalgenome.org/).

The Genome Research on Atlantic Salmon Project (GRASP) has been conducted in Canada, where Atlantic salmon (*Salmo salar*) is an important farmed fish. In this project, genetic linkage maps and physical maps have been constructed for the Atlantic salmon genome. Genome reagents and tools have been prepared, including large numbers of expressed sequence tags (ESTs), bacterial artificial chromosome (BAC) end sequences (BES), and microarray platforms. This project has been renewed and re-termed as the cGRASP project (Consortium for Genomic Research on All Salmonids Project; http://web.uvic.ca/cbr/grasp/).

Several major genome projects have been conducted and initiated in Europe, including the SalMap project for mapping the Atlantic salmon genome, the BASSMAP project for mapping the genome of the European sea bass (*Dicentrarchus labrax*), the BRIDGEMAP project for genome of sea bream (*Sparus aurata*), and a metagenomics project focused on evolution of genome components coping with environmental stresses. The BRIDGEMAP project is a collaborative project funded by the EU initiated in 2001 with three major objectives: 1) construction of a framework genetic linkage map of sea bream for aquaculture as well as for population monitoring for fisheries; 2) To develop basic genome tools and reagents from sea bream for use in comparative genomics across species; and 3) Construction of physical maps using
bacterial artificial chromosome (BAC) libraries and radiation hybrid mapping panels (http://www.bridgemap.tuc.gr/).

In China, the Grass Carp Genome Project was initiated in 2005 and officially announced in the 13th International Congress On Genes, Gene Families And Isozymes (http://www.cafs.ac.cn/page/cafs/guanggao/jiyin/show1eng.htm). This project involves multiple institutions in China and is aimed at producing genome reagents like ESTs, BAC libraries, BAC end sequences, physical maps, linkage maps, before eventually sequencing the entire genome of the grass carp (Ctenopharyngodon idella). Genome studies of shrimps have been conducted for several years in China and Thailand and ESTs and microarrays have been produced; linkage maps have also been constructed (http://pmonodon.biotec.or.th/; Wang et al., 2006). In Japan, genome projects have been conducted with Japanese flounder, yellow tail, shrimps, and oysters. Fish genome studies are in progress in many parts of the world, but the major efforts are located in the United States of America, Canada, Europe, China, and Japan. Genome projects are expensive and many developing countries cannot afford them.

Major achievements of aquaculture genomics
Framework genetic linkage maps have been established in salmon, trout, tilapia, catfish, shrimp, oysters, and many other species (Table 1). Large numbers of molecular markers have been developed and efforts for mapping more markers are increasing. Basic genome reagents have been or are now being established for farmed finfish, crustaceans and molluscs. Large-insert DNA libraries, such as BAC libraries, are available for Atlantic salmon, rainbow trout (Oncorhynchus mykiss), tilapia (Oreochromis spp.), channel catfish (Ictalurus punctatus), and several other finfish species. Two BAC libraries have been constructed for oysters and those for shrimps are being constructed. Physical maps have been constructed in Atlantic salmon (Ng et al., 2005), Nile tilapia (Oreochromis niloticus) (Katagiri et al., 2005), and channel catfish (Xu et al., 2007). Gene discovery efforts through sequencing ESTs are increasing. A total of almost one million ESTs are now available from farmed aquatic species, of which a large percentage is from Atlantic salmon, rainbow trout, and channel catfish. The Joint Genome Institute (JGI) of the US Department of Energy (DOE) has initiated large EST projects for channel catfish (to produce 600,000 ESTs, John Liu of Auburn University serves as the principal investigator), oysters (to produce 600,000 ESTs, Dennis Hedgcock of the University of Southern California serves as the principal investigator), and genome survey project in tilapia-related species (to sequence a total of 10% genome coverage from five tilapia-related species, Thomas Kocher of the University of New Hampshire serves as the principal investigator). Descriptions of these JGI sequencing projects can be found at http://www.jgi.doe.gov/News/news_5_12_05.html. cDNA microarray technologies have been developed and used in Atlantic salmon, shrimps, oysters, and channel catfish. Although farmed fish genome research had a late start, this allowed researchers to learn lessons from scientists working with other species, and more advanced genome technology also provided greater efficiency.

Research on the genomes of farmed fish has focused on performance and production traits such as growth rate, feed conversion efficiency, disease resistance, tolerance to environmental stresses such as high ammonia, low dissolved oxygen, tolerance to cold temperature and to various salinities. In most cases, the genomes of farmed aquatic species are smaller than, or comparable to the human genome. Many farmed fish species have high fecundity, which provides opportunities to create large resource and reference families that allow great selection pressure to be applied at the phenotypic level for the analysis of quantitative trait loci (QTL). Experiments can be repeated many times as the related expense is relatively small. The large size of resource families allows accurate mapping of important genes responsible for traits. However,
<table>
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<th>Marker system used</th>
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<td>AFLP</td>
<td>Li et al., 2003</td>
</tr>
<tr>
<td><em>Penaeus chinensis</em></td>
<td>Chinese shrimp</td>
<td>AFLP</td>
<td>Li et al., 2006</td>
</tr>
<tr>
<td><strong>Molluscs</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Crassostrea virginica</em></td>
<td>Eastern oyster</td>
<td>Microsatellites</td>
<td>Yu and Guo, 2003</td>
</tr>
<tr>
<td><em>Crassostrea gigas</em></td>
<td>Pacific oyster</td>
<td>AFLP, Microsatellites</td>
<td>Li and Guo, 2004; Hubert and Hedgecock, 2004</td>
</tr>
<tr>
<td><em>Chlamys farreri</em></td>
<td>Zhikong scallop</td>
<td>AFLP</td>
<td>Li et al., 2005</td>
</tr>
<tr>
<td><em>Haliotis discus hannae</em></td>
<td>Pacific abalone</td>
<td>AFLP, RAPD, Microsatellites</td>
<td>Liu et al., 2006; Sekino and Hara, 2007</td>
</tr>
<tr>
<td><strong>Other</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Stongylocentrotus nudus</strong></td>
<td>Sea urchin</td>
<td>AFLP</td>
<td>Zhou et al., 2006</td>
</tr>
</tbody>
</table>
farmed fish genomics faces great challenges because there are many farmed fish species and funding levels for aquaculture genomics have been low. Technically, labeling of individual fish in research is often a daunting task, unless intrusive procedures are used; the aquatic environment also makes real time observations extremely difficult; genome duplication is widespread in teleost fishes, which poses challenges related to complexities for the analysis of gene arrangement, expression and function.

**Sequencing farmed fish genomes**

In spite of the large effort in genome sequencing of the zebrafish (*Danio rerio*) (http://www.sanger.ac.uk/Projects/D_rerio/), green spotted pufferfish (*Tetraodon nigroviridis*) (http://www.genoscope.cns.fr/externe/English/Projets/Projet_C/), Japanese pufferfish (*Fugu rubripes*) (http://www.genomenewsnetwork.org/articles/1101/Pufferfish_sequenced.shtml) and medaka (*Oryzias latipes*) (http://dolphin.lab.nig.ac.jp/medaka/), no genomes of aquaculture species have been sequenced. White papers have been generated advocating sequencing a few genomes of important aquaculture species including those for rainbow trout (http://www.animalgenome.org/aquaculture/salmonids/RainbowProposal.pdf) and oysters (http://findarticles.com/p/articles/mi_m0QPU/is_2_24/ai_n15390229). The major constraints for sequencing the fish genomes are financial. In the overall genomics revolution, there has been little attention so far to the genomics of farmed fish, even for the most important species. Plant and livestock genomes have been given priority. Of the major agricultural animals, bovine, swine, and chicken genomes are being sequenced. The entire genome sequences would provide research and application advantages for the involved species. Taking USDA funding priorities as an example, only species whose genome is sequenced with a draft sequence over five fold of the genome coverage (i.e., the total base pairs sequenced should be greater than five times of the genome size of the species) is eligible for funding from the Functional Genomics Program. Obviously, no species used in aquaculture meets the criteria. All aquaculture species are thus currently excluded from its funding. However, emerging sequencing technologies might enable genomes to be sequenced more cheaply and efficiently and then it should be possible to sequence important farmed fish genomes. With or without this, fish genomics should focus on FIGR management (use and conservation), genetic enhancement, and the environmental aspects of fish genomes. Among these, genetic enhancement often takes the priority, but it is very important to note that FIGR management, and environmental genetic issues are often directly related to genetic enhancement, and should be given much greater attention.

5. ACTUAL AND POTENTIAL APPLICATIONS OF GENOMICS AND ANALYTICAL GENETIC TECHNOLOGIES IN FISH GENETIC RESOURCES MANAGEMENT

Aquaculture production is growing rapidly to provide food fish for the world’s rapidly growing population and now provides approximately 40% of food fish consumed by humans (FAO, 2006). Many capture fisheries are currently harvested at or above maximum sustainable levels, and are in global decline because of over-harvesting and habitat degradation. Wild fish genetic resources (FiGR) are being depleted and some are facing extinction. Some aquaculture operations still depend on wild FiGR for seed and broodstock and wild fish populations are important resources for fish genetics research, including breeding programs and genomics.

The genomics revolution and genetic analytical technologies have many actual and potential applications for capture fisheries and aquaculture, including FiGR management. Their practical applications in aquaculture include, *inter alia*, marker-assisted selection (MAS), environment protection, genetic management of broodstocks, and genetic improvement of framed fish. Analytical genetic technologies will contribute
to lessening the adverse environmental impacts of aquaculture as well as to resolution of the disease problems through genetic enhancement, development of effective vaccines and their delivery systems, and development of rapid and accurate diagnostic tools. Future applications also include the safe use of transgenic technologies. Annex 1 summarizes major genomics and other methods of genetic analysis for application to natural and farmed aquatic species.

Diseases are major threats to sustainable aquaculture, in crustacean farming, especially shrimp farming, and in the farming of some molluscs, especially oysters. Diseases also affect the farming of many types of finfish, including carps, catfish and salmonids. Countermeasures to ensure the health and survival of farmed fish, including genetic technologies, are much needed. Superior broodstocks resistant to major diseases are needed. Although rich genetic resources must exist among aquaculture species for resistance to major fish diseases, for fast growth and for efficient feed conversion, genomic research is required to identify and then utilize these. Resistance-linked markers are especially needed for marker-assisted selection. Direct selection of disease resistance has proven to be very difficult in aquaculture. Genome based-technologies could provide solutions to meet some of the challenges presented by economically important pathogens. Genetic technologies for increasing and identifying disease resistance have proven safe, reliable, and environmentally sound for livestock. Mapping of large numbers of markers will pave the way for seeking QTLs for disease resistance in fish. This will add a new dimension to the new generation of technology for genetic improvement of disease resistance through marker-assisted selection in aquaculture. Genome research should facilitate marker-assisted selection for genetic improvement in many production traits of farmed fish. Some recent QTL studies are listed in Table 2.

**Genetic improvement through marker-assisted selection**

Marker-assisted selection is a major potential application that is used as an argument for expanding research on the genomics of farmed fish. DNA marker technologies are already used routinely for stock identification is routine in some farmed fish species (Beacham et al., 2000; 2005, Duchesne and Bernatchez, 2007). A few markers linked with performance and production traits have been identified (Moen et al., 2004a; 2004b; Martyniuk et al., 2003; O’Malley et al., 2003; Reid et al., 2005; Nichols et al., 2003a; Palti et al., 2001; Cnaani et al., 2003). However, despite a great deal of promise, only a handful of cases demonstrating practical usefulness of MAS in reducing frequencies of recessive alleles causing genetic diseases, determining simple Mendelian traits and improvement in a few species have been reported so far (Dentine, 1999; Dekkers, 2004). In livestock, commercial implementation of MAS related to improvement of quantitative traits has been employed for removal of deleterious major genes, growth rate, meat quality, disease resistance and reproductive traits in pigs and in other species such as cattle where markers are used routinely for improvement of protein percentage in milk and marbling and tenderness in beef cattle. Dekkers (2004) has summarized the progress and use of markers as “the current attitude toward MAS is … cautious optimism”. Fine QTL mapping will allow traits-linked markers to be identified and used for MAS, which should not replace traditional selective breeding, but should complement to ensure accurate and effective selection and to contribute indicators at the molecular level that phenotypically selected fish truly contain the genes that breeders believe they do. Specific studies related to performance and production traits for aquaculture species must be accomplished before applying MAS in aquaculture. With low levels of funding, it is anticipated that such research may still requires years of effort, and actual MAS in aquaculture may still need at least five years.
Protection of the environment, with increasing aquaculture production and profitability

Aquaculture has a great challenge ahead to address potential environmental impacts and genomics can make significant contributions. Genome research should focus on how to benefit aquaculture and at the same time on how to protect the environment. Through research on the genomes of farmed fish, new technologies can be developed for monitoring the aquatic environment using bioindicators, biomarkers and genome expression signatures (e.g. Almeida et al., 2005; Gracey and Cossins, 2003; Cossins and Crawford, 2005). Environmental genomics is therefore an important focus for farmed fish genomics. Metagenomics and ecogenomics include the goal of using genome technologies to improve environmental quality. Environmental genomics is now a major driving force (e.g. Travis et al., 2003; MacGregor, 2003; Frazier et al., 2003; Almeida et al., 2005).
**Disease diagnosis, food safety, disease resistance, fish vaccines, drug-resistant pathogens**

Genomics can contribute much to the accurate diagnosis of fish diseases and to ensuring the safety of aquatic produce. Existing technologies are practical and capable of delivering results immediately (Kerr and Cunningham, 2006; Adams and Thompson, 2006).

Genome research, through QTL mapping, MAS and transgenesis, provides potential avenues for addressing some of the disease problems that threaten aquaculture. Through QTL mapping, it is possible to locate major genes responsible for disease resistance. Through MAS, brood stocks can be developed containing disease resistance genes (e.g. Palti et al., 1999; Moen et al., 2004b; Nichols et al., 2003a) and conceptually, traditional hybridization can be used to allow their introgression (Liu et al., 2003; Senanan et al., 2004).

Vaccines should be developed for fish as for livestock, though their applications in aquaculture have some limitations (Lorenzen et al., 2002; Evensen et al., 2005). One major difference is the large number of individuals in populations of farmed fish species and their relatively low individual value. The aquatic environment also poses technical difficulties. Genome research may allow development of more effective vaccines, including DNA vaccines (Kurath, 2005). Effective vaccine delivery systems must also be developed. Genome technologies should also provide means for monitoring drug resistance in fish pathogens. This is a significant problem because countries that produce fish but have relaxed laws with respect to drug use in aquaculture could contribute to increases in drug resistance in many pathogens, including some that affect humans (Graslund and Bengtsson, 2001; Cabello, 2006).

**Genetic characterization**

Genetic marking and identification of fish species, strains, lines, populations, and individuals is very important not only for aquaculture and hatchery operations, but also for capture fisheries management. Genome technologies have the capacity to provide “diagnostic kits” to identify many important species and populations, using DNA marker technologies (for reviews, see Liu and Cordes, 2004; Grant, 2007; Pullin, 2007; Smith, 2007).

DNA marker technologies should be adapted for wider use in the characterization of wild FiGR. This is particularly important for fish species and stocks that are captured by humans, because some fisheries might be depleting FiGR. For endangered species and stocks, genome technologies can characterize fish produce even after it has been cooked. This should provide greater levels of law enforcement. Consumers should also be protected to ensure that fish produce is labeled accurately at the point of sale (Maldini et al., 2006).

**Transgenic fish**

Early attempts to develop transgenic fish were hindered by a lack of fish promoters and much of the early research was conducted with viral promoters (Dunham and Liu, 2006). Gene-based genetic improvements have now been well demonstrated in fish species using transgenic technologies. In spite of low public acceptance, transgenic work in salmon has demonstrated that growth rate can be enhanced over 10 times by transferring only a growth hormone gene (Du et al., 1992; Roberts et al., 2004; Devlin et al., 2004), illustrating the plasticity of some fish genomes and their functions. Other transgenic fish have been developed with improved growth rate, color, disease resistance, survival in cold and body composition, and the ability to produce pharmaceutical proteins. Transgenic zebrafish with altered coloration have been commercialized and
applications are pending for commercialization of transgenic salmon, carp and tilapia transgenes with transferred growth hormone genes (for examples of reviews, see Devlin et al., 2006; Kapuscinski, 2005; Domergue et al., 2005; Fu et al., 2005; Napier et al., 2004; Zbikowska, 2003; FAO, 2000; Maclean, 2000; Zhu and Sun, 2000; Iyengar et al., 1996; Chen et al., 1996; Gong and Hew, 1995; Hew et al., 1992; Houdebine and Chourrout, 1991; Chen and Powers, 1990). To minimize environmental risks, additional technologies such as transgenic sterilization need to be developed (Dunham and Liu, 2006). Genomic research has produced an abundance of molecular genetic information including many genes for consideration for gene transfer, highly regulated gene promoters, and knowledge about their expression and function. Functional genomics analysis should be applied in the future to enhance the capacity and versatility of transgenic technology, and to facilitate assessment of the biosafety aspects of development and use of transgenic fish. Increased research will be needed for determining environmental risk, measuring the fitness of transgenic fish and for determining the safety of aquatic produce derived from them. The future success and application of transgenic fish will be dictated by successful demonstration of acceptable environmental risk, assurance of food safety, appropriate government regulation and labeling, public education and opinion, and development of genetic sterilization for transgenic fish. Where commercial production of transgenic food fish is the objective, fish promoters should be used. Advances in genomics will provide these as well as important genes for gene transfer that could have greater public acceptance.

Some important commercial traits of farmed fish - such as resistance to diseases, feed conversion efficiency, tolerance to poor water quality, harvestability, carcass yield, increased reproduction and improved utilization of plant resources have yet to be addressed by transgenic technology. Basic information from genomic research may be the starting point to address effectively genetic enhancement of these traits. One of the greatest future potential benefits of gene transfer in fish could be enhancement of disease resistance in fish. Transgenic fish with enhanced disease resistance would increase profitability, production, efficiency and the welfare of the cultured fish. Preliminary research (Dunham et al., 2002; Chiou et al., 2002; Sarmasik et al., 2002) indicates great promise for success of this approach for enhancing disease resistance. The use of transgenic fish in recreational fisheries could involve release of transgenic fish into open waters or into more confined, urban environments. Public opinion will vary in regards to this application and the use of transgenic fish in aquaculture of food fish and ornamental fish will likely occur much earlier than their use in recreational fisheries. In the ornamental fish trade, a transgenic petfish named Glofish has already been marketed (Gong et al., 2002, 2003).

**Combining genetic technologies**

Transgenic technology is no silver bullet; neither are genome technologies or traditional selective breeding. Genomics and combined genetic technologies are expected to lead to a much larger scope of genetically improved farmed fish. Partly, this is because the history of domestication and selective breeding of many farmed fish species has been short and great potentials for genetic improvement have yet to be realized. Continued selection plus the application of MAS will likely overcome many of the challenges faced by traditional selection alone, and provide faster and more effective results. Markers for complex traits are more difficult to be identified, and usually complex traits are controlled by multiple genes. Nonetheless, MAS has great potential to fulfill the promises made by agricultural genomics.
6. ENVIRONMENTAL AND SOCIAL ISSUES

A number of governmental and non-governmental organizations (NGO) have started discussions on issues of genomics related to ethics, environment, economy, law, and society (GE3LS). Genome Canada (http://www.genomecanada.ca/) has conducted annual GE3LS symposium for several years focusing on conflicting worldviews, social cohesion, ownership, and the democratic deficit. These themes were explored in relation to the application of genomics and proteomics to the fields of agriculture, environment, fisheries and forestry. In the United States of America, the Department of Energy (DOE) and the National Institutes of Health (NIH) Genome Programs set aside 3% to 5% of their respective annual Human Genome Project budgets for the study of the project’s ethical, legal, and social issues (ELSI) (http://www.ornl.gov/sci/techresources/Human_Genome/elsi/elsi.shtml). Many of the issues and concerns discussed in this section were obtained from this website because published papers are scarce in this area. As the GE3LS issues started to emerge with human genome related issues, many of the similar concerns related to genomics will emerge in aquaculture and fisheries related areas.

GMOs in aquaculture

An important issue in aquaculture and fisheries is the application of GMOs, the genetically modified organisms produced through the use of genetic engineering. Other issues include genetic impact of farmed and ornamental fish on wild populations, the ownership of and access to FiGR, and imbalances of genome technologies and capacities in various parts of the world.

The linkage of genomics to biotechnology and transgenic technology is its ability to rapidly discover, identify and characterize genes of economic importance. Such genes can be used for biotechnology, pharmaceutical purposes, or transgenics. A number of controversies exist concerning the use of genetically engineered organisms. The supporting forces come from the benefits GMO’s can bring to the society. The fundamental argument for the development of GMOs is the increased food production efficiency for growing human populations.

Researchers at the University of Guelph have developed a new breed of Yorkshire pigs trademarked Enviropig™ that use plant phosphorus more efficiency (Golovan et al., 2001a, 2001b). Non-transgenic pigs are unable to use an indigestible form of phosphorus called phytate present in the cereal grain diet. Therefore producers add supplemental phosphate or phytase enzyme to the diet in order to meet the phosphorus requirement for optimal growth and development. The novel trait of the Enviropig™ enables it to degrade the indigestible phytate and absorb the phosphate eliminating the need to supplement the diet with readily available phosphate, and as a consequence the phosphorus content of the manure is reduced by as much as 60%. Digestion of the phytate also leads to improvements in digestion of minerals in the diet. Clearly, potential use of such transgenic animals would have positive impact to the environment, and such research may inspire similar research in fish.

Consumer choice

Informing consumers of fish products about their identity and origin, by accurate labeling, facilitates consumer choice and product development and marketing. However, economic interests and international trade politics may seriously constrain product labeling. Labeling is not yet mandatory in some countries (e.g. United States of America). Another consideration is consumer awareness or public education. A survey conducted in New Jersey (http://www.nal.usda.gov/bic/Pubpercep/) found that most residents (91%) felt they had an “adequate” or “very good” understanding of how food is grown and produced. However, much of the public was unfamiliar
TABLE 3

Existing genomic reagents and tools for important aquaculture and fisheries species. All information was obtained from the National Center for Biotechnology Information (NCBI) (http://www.ncbi.nlm.nih.gov/) by using Entrez, the Life Sciences Search Engine as of 6 December 2006. Expressed sequence tags represent sequence reads from single pass sequencing of transcribed sequences, while total DNA sequences are number of sequence reads obtained from both expressed sequences and genomic sequences.

<table>
<thead>
<tr>
<th>Common name</th>
<th>Scientific name</th>
<th>Importance in aquaculture, capture Fisheries, or both</th>
<th>Number of expressed sequence tags (ESTs)</th>
<th>Total DNA sequences</th>
<th>Characterized proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atlantic salmon</td>
<td>Salmo salar</td>
<td>Both</td>
<td>430 340</td>
<td>434 380</td>
<td>1 380</td>
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<tr>
<td>Rainbow trout</td>
<td>Oncorhynchus mykiss</td>
<td>Aquaculture</td>
<td>262 256</td>
<td>265 613</td>
<td>2 727</td>
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<td>Catfish</td>
<td>Ictalurus spp.</td>
<td>Aquaculture</td>
<td>57 084</td>
<td>79 108</td>
<td>1 641</td>
</tr>
<tr>
<td>Common carp</td>
<td>Cyprinus carpio</td>
<td>Aquaculture</td>
<td>19 344</td>
<td>20 555</td>
<td>1 099</td>
</tr>
<tr>
<td>Eastern oyster</td>
<td>Crassostrea virginica</td>
<td>Aquaculture</td>
<td>9 018</td>
<td>9 125</td>
<td>73</td>
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<tr>
<td>Tilapia</td>
<td>Oreoichromis spp.</td>
<td>Both</td>
<td>676</td>
<td>6 688</td>
<td>519</td>
</tr>
<tr>
<td>Pacific oyster</td>
<td>Crassostrea gigas</td>
<td>Both</td>
<td>4 201</td>
<td>5 259</td>
<td>284</td>
</tr>
<tr>
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<td>Chlamys farreri</td>
<td>Both</td>
<td>3 466</td>
<td>3 598</td>
<td>208</td>
</tr>
<tr>
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<td>1 055</td>
<td>1 055</td>
<td>201</td>
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<td>534</td>
<td>809</td>
<td>227</td>
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<td>Skipjack tuna</td>
<td>Katsuwonus pelamis</td>
<td>Fisheries</td>
<td>0</td>
<td>323</td>
<td>77</td>
</tr>
<tr>
<td>Japanese anchovy</td>
<td>Engraulis japonicus</td>
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<td>238</td>
<td>127</td>
</tr>
<tr>
<td>Alaska Pollock</td>
<td>Theragra chalcogramma</td>
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<td>161</td>
<td>252</td>
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<tr>
<td>Silver carp</td>
<td>Hypophthalmichthys molitrix</td>
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<td>84</td>
<td>19</td>
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<tr>
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<td>48</td>
<td>10</td>
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<td>Fisheries</td>
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<td>Fisheries</td>
<td>0</td>
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<td>Trichiurus lepturus</td>
<td>Fisheries</td>
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<td>0</td>
</tr>
</tbody>
</table>

with traditional methods for producing hybrid plants and animals. While over half the population (54%) acknowledged that they had heard of cross-fertilization or cross-breeding, only 28 percent said they had eaten a fruit or vegetable produced by this method. This is surprising because most commercially available fruit and vegetables are now hybrids. Even more surprising, 17% of the population interviewed believed that they have eaten a fruit or vegetable produced by genetic engineering, though no such product is yet available. This demonstrates the great need for improving public education and awareness about applications of genetics in food production.

Geographical distribution of fish production and fish genomics research
Developed countries will likely play leading roles in the development of farmed fish genomics and genetic technologies that in turn will enhance aquaculture production. However, most of the world’s farmed fish production, comes from developing countries...
How the farmed fish genome technologies will be disseminated is a major concern. Efforts should be made to promote international cooperation and collaboration in genome research and utilization of results and products.

Genetic maps have yet to be developed for many important farmed fish and for many important species that are targeted by capture fisheries. Some of the available genome reagents and resources of fish genomic research so far are summarized in Table 3, from which it is clear that the major genome research activities have been focused on farmed fish species. Much more international collaboration is needed in order to enhance genomics research efforts on major farmed fish and capture fisheries species of the world. Microsatellites are needed for genetic linkage mapping and mapping of QTL, as well as for population genetic studies. ESTs are needed for analysis of gene expression, and also ESTs serve as rich sources for polymorphic markers, and serve as material basis for the development of microarrays unless otherwise the genome sequence is available. Genome sequence surveys (GSS) allow sampling of the genome for the assessment of the genome composition, repeat structure, as well as for polymorphic marker identification.

Need for globally accessible information
There are currently no comprehensive globally accessible databases on FiGR. It will be important to gather information on intraspecific genetic diversity for major capture fisheries species and major farmed fish species (Pullin, 2007). Databases of DNA fingerprints will help in species identification and also in the interest of protection of endangered species and the consumer’s interest (Smith, 2007). Law enforcement agencies have trouble in identification of fish in markets and served in restaurants. In many cases, endangered species are involved, but unless more effective genome technologies are developed to provide rapid and accurate identification using fingerprinting techniques, it is difficult to provide effective means for the protection of endangered species. In some cases, inferior fish products are mislabeled as having come from more expensive species. Genome research on major farmed fish species has generated molecular markers allowing population studies and genetic resource analysis. In contrast, little genome information exists for most capture fisheries species and this is limiting the application of genome technologies in assessment of the status and conservation of wild FiGR.

Public education
The public is generally ill-informed and naïve about biotechnology in food production, including the pros and cons of transgenics, genomics and genetic technologies. Public education should be considered an important issue. Many professionals in capture fisheries and aquaculture also do not understand well the potentials and implications of genomics and genetic technologies. While information dissemination from genomics to the public is very important, better exchanges of information between genome researchers and aquaculture and fisheries professionals are also essential. As noted by Dr. Alex Mackenzie, Vice President of Research at Genome Canada, “in order for us to conduct ourselves optimally as a society, I think an informed populace is our most potent weapon” (http://www.iog.ca/about_us.asp?pageid=28).

7. REFERENCES


with 2 other species of fish (Arctic charr and Atlantic salmon) within the tetraploid derivative Salmonidae family (subfamily: *Salmoninae*). *Genome*, 48: 1037-1051.


profiling following DNA vaccination of rainbow trout against infectious hematopoietic necrosis virus. *Molecular Immunology*, 43(13): 2089-2106.


ANNEX 1

Genetic marker, mapping and other technologies

Allozyme markers
Allozymes”, or “allelic isozymes”, are the different allelic forms of the same enzymes encoded at the same locus (Parker et al., 1998). The most common use of allozyme electrophoresis is to detect genetic variation in natural populations. In the last 30 years, large amounts of allelic frequency data were collected from many fish species for management purposes. Although use of allozyme data in aquaculture appears to be limited compared to its use in capture fisheries population studies, aquaculture has utilized this information for its development because aquaculture and fisheries sometimes cannot be separated from each other (Dunham 2004). Allozyme electrophoresis in aquaculture is used for stock identification, parentage analysis, hybrid identification, inbreeding analysis and limited genetic mapping (Liu and Cordes 2004). Although allozyme studies have not found common application in marker assisted selection, correlations between certain allozyme markers and performance traits has been reported (Hallerman et al., 1986). Similarly, due to the limited number of polymorphic loci available, use of these markers in linkage mapping in fish is limited.

The major drawback of allozyme analysis is the necessity for a large amount of fresh or frozen tissue samples. This often requires lethal sampling, especially if a full array of allozyme markers is to be studied. Furthermore, although allozymes represent actual gene products, they often measure a very small portion of the genomic variation because a limited number of loci are involved (Utter et al., 1987). Although cheap and technically easy, numbers of allozyme loci and polymorphisms are low (Agnèse et al., 1997). These drawbacks seriously limit the applications of allozymes for genome studies.

Mitochondrial DNA markers
The mitochondrial genome evolves more rapidly than the nuclear genome. The rapid evolution of the mtDNA makes it highly polymorphic within a given species. Mitochondrial DNA is maternally inherited for the most part, but there are reports of paternal leakage during fertilization (Birky et al., 1989). Mitochondrial DNA analysis is actually a restriction fragment length polymorphism (RFLP) analysis except that the target molecule is mtDNA rather than nuclear genomic DNA (Liu and Cordes 2004). The high levels of polymorphism, the maternal inheritance and the relatively small size of mtDNA make the RFLP analysis using mtDNA one of the easiest methods for many population studies (Okumus and Ciftci, 2003; Liu and Cordes, 2004, May, 2003; Billington, 2003). Mitochondrial DNA markers have been used extensively to analyze genetic variation in several different aquaculture species including striped bass (Wirgin et al., 1991; Garber and Sullivan, 2006), channel catfish (Waldbieser et al., 2003), walleye (Merker and Woodroff, 1996), salmonids (Nilesen et al., 1998, Crespi and Fulton 2004), red snapper (Pruett et al., 2005), and bluegill (Chapman, 1989). There are two major drawbacks of mtDNA markers. One is the non-Mendelian inheritance of mtDNA; and the second is the proportion of the total genomic variation one can observe with mtDNA alone. These characteristics place limitations to the validity of using mtDNA for genetic studies.

RFLP markers
Restriction fragment length polymorphism (RFLP) was the most popular approach for analysis of genetic variation during the 1980's. As indicated by its name, RFLP is based on DNA fragment length differences after digesting genomic DNA with one or more restriction enzymes. In spite of its earlier popularity, RFLP is able to
detect only large shifts in DNA fragment sizes. It is unable to detect the vast majority of point mutations. As a result, polymorphic rates are low at most loci. The efforts involved in RFLP marker development have been enormous. RFLP attempts to detect genetic variation one locus at a time. The low polymorphic rates, when coupled with expensive and laborious processes, have made application of RFLP limited. It should be particularly noted that RFLP requires previous genetic information, such as the availability of probes or sequence information, information often not available for many fish or other aquaculture species. Future use of RFLP will focus on analysis of Single Nucleotide Polymorphisms (SNP) residing within restriction sites.

**RAPD markers**

Random amplified polymorphic DNA (RAPD) is a PCR-based multilocus DNA fingerprinting technique (Welsh and McClelland, 1990; Williams et al., 1990). RAPD markers are inherited as Mendelian markers in a dominant fashion. RAPDs have all the advantages of a PCR-based marker, with the added benefit that primers are commercially available and do not require prior knowledge of the target DNA sequence or gene organization. Other advantages of RAPDs are the ease with which a large number of loci and individuals can be screened. The major weakness of RAPD is its low reproducibility due to the use of low annealing temperatures, and its dominant mode of inheritance. RAPD markers have been widely used for species and strain identification in fishes (Partis and Wells, 1996; Liu et al., 1998; 1999) and mollusks (Klinbunga et al., 2000; Crossland et al., 1993), analysis of population structure in black tiger shrimp (Tassanakajon et al., 1998) and marine algae (Van Oppen et al., 1996), analysis of genetic impact of environmental stressors (Bagley et al., 2001), and analysis of genetic diversity (Wolfus et al., 1997; Hirschfeld et al., 1999; Yue et al., 2002). RAPD markers have also been used for linkage mapping in fish species (Table 1). However, as more efficient and reliable marker systems such as AFLP emerged, the use of RAPD markers in genome research declined rapidly. However, it is a very useful marker system for rapid hybrid identification, strain identification, and population studies in fisheries species where other genomic information may be lacking.

**AFLP markers**

Amplified fragment length polymorphism (AFLP) is a PCR-based DNA fingerprinting technique that provides robust analysis of the genome variations. AFLP markers are inherited in a Mendelian fashion as dominant markers. Several major strengths make AFLP markers of choice in many situations. First AFLP requires no prior molecular information for application to the species of interest. This is particularly useful for aquaculture and fisheries species where molecular information is often not available. Second, AFLP is highly robust allowing generation of a large number of polymorphic markers with limited efforts and resources. Third, it is highly reproducible and reliable. The major weakness of AFLP markers is their dominant nature of inheritance. AFLP is more technically demanding, requiring special equipment such as automated DNA sequencers for optimal operations. AFLP is well adapted for many types of genetic analysis such as analysis of genetic diversity, population structures, migration, hybrid identification, strain identification, parentage identification, genetic resource analysis, reproduction contribution, endangered species protection, marker-assisted selection, and genome mapping. Despite the advantages of AFLP, published literature on its application for the analysis of genetic variation of fish population genetic studies is still limited due to technical difficulties and requirement for special equipment (Seki et al., 1999; Jorde et al., 1999; Sun et al., 1999; Chong et al., 2000; Kai et al., 2002; Mickett et al., 2003; Whitehead et al., 2003; Mock et al., 2004; Campbell and Bernatchez, 2004; Simmons et al., 2006). Many AFLP analyses in fish so far have been limited to genetic linkage analysis (Table 1), and analysis of parental genetic contribution involving
interspecific hybridization (Youngson et al., 2001) and meiogynogenesis (Felip et al., 2005). In a recent study of the black rockfish (*Sebastes inermis*), Kai et al. (2002) used AFLP to distinguish three color morphotypes, in which diagnostic AFLP loci were identified as well as loci with significant frequency differences. In such reproductive isolated populations, it is likely that “fixed markers” of AFLP can be identified to serve as diagnostic markers. Fixed markers are associated most often with relatively less migratory, reproductive isolated populations. With highly migratory fish species, fixed markers may not be available. However, distinct populations are readily differentiated by difference in allele frequencies. For instance, Chong et al. (2000) used AFLP for the analysis of five geographical populations of Malaysian river catfish (*Mystus nemurus*) and found that AFLP was more efficient for the differentiation of sub-populations and for the identification of genotypes within the populations than RAPD although similar clusters of the populations were concluded with either analysis. Genetic resource diversity have been assessed using AFLP (Micketti, 2003), and a comparison of the aquacultured catfish with wild populations suggested that the domestic fish had much narrower genetic diversity (Simmons et al., 2006). The impact of the aquaculture catfish on wild catfish populations was found to be little, if any (Simmons et al., 2006).

**Microsatellite markers**

Microsatellites are tandemly arranged simple sequence repeats (Tautz and Renz, 1984; 1989). Microsatellites are highly abundant in various eukaryotic genomes including all aquaculture species studied to date. Generally speaking, more compact genomes tend to contain smaller proportion of repeats including simple sequence repeats. For example, the highly compact genome of Japanese pufferfish contains 1.29% microsatellites (Crollius et al., 2000). During a genomic sequencing survey of channel catfish, microsatellites were found to represent 2.58% of the catfish genome (Xu et al., 2006). In most fish species, dinucleotide (AC) repeats are the most abundant forms of microsatellites. Microsatellites are highly polymorphic such that they are suitable for differentiation of individuals, as well as populations, and species. Microsatellites are inherited in a Mendelian fashion as co-dominant markers. As microsatellites have the greatest differentiating power, they have been widely used in aquaculture and fisheries in areas including linkage mapping (Table 1, Liu and Cordes, 2004; Chistiakov et al., 2006), analysis of genetic diversity, population genetics and conservation genetic analysis, parentage analysis, molecular epidemiology and pathology, QTL mapping (Chistiakov et al., 2006). Microsatellites are highly adaptable for marker-assisted selection, but have not been applied in aquaculture yet because the linkage maps and QTL analysis for important traits are still lacking.

**SNP markers**

Single nucleotide polymorphisms (SNPs) are alternative bases at a given nucleotide position. Such sequence differences due to base substitutions have been well characterized since the beginning of DNA sequencing in 1977, but the ability to genotype SNPs rapidly in large numbers of samples was not possible until in the late 1990s. SNPs are becoming a focal point in molecular marker development since they are the most abundant polymorphism in any organism, adaptable to automation, and reveal hidden polymorphism not detected with other markers and methods. Theoretically, a SNP within a locus can produce as many as four alleles, however, most SNPs are usually restricted to one of two alleles and have been regarded as bi-allelic. SNP markers are inherited as co-dominant markers. Several approaches have been used for SNP discovery including SSCP analysis (Hecker *et al*., 1999), heteroduplex analysis (Sorrentino *et al*., 1992), and direct DNA sequencing. DNA sequencing has been the most accurate and most-used approach for SNP discovery. Random shotgun
sequencing, amplicon sequencing using PCR, and comparative EST analysis are among the most popular sequencing methods for SNP discovery.

Despite technological advances, SNP genotyping is still a challenging endeavor and requires specialized equipment. Traditional methods available for SNP genotyping include: direct sequencing, single base sequencing (reviewed by Cotton, 1993), allele-specific oligonucleotide (ASO, Malmgren et al., 1996), denaturing gradient gel electrophoresis (DGGE, Cariello et al., 1988), single strand conformational polymorphism assays (SSCP, Suzuki et al., 1990), and ligation chain reaction (LCR, Kalin et al., 1992). Each approach has its advantages and limitations, but all are still useful for SNP genotyping, especially in small laboratories limited by budget and labor constraints. Large-scale analysis of SNP markers, however, depends on the availability of expensive, cutting-edge equipment. Several options are available for efficient genotyping using state of the art equipment. Particularly popular are methods involving MALDI-TOF (Matrix-assisted laser desorption ionization - time of flight) mass spectrometry (Ross et al., 1998; Storm et al., 2003), pyrosequencing (Ahmadian et al., 2000; Alderborn et al., 2000; He et al., 2003), Taqman allelic discrimination (Li et al., 2004), real-time (quantitative) PCR (Nurmi et al., 2001), and the use of microarray or gene chips (Hacia et al., 1999). Mass spectrometry and microarray technologies require a large investment in equipment. The equipment for pyrosequencing and quantitative PCR is generally under $100,000, and should be more affordable to many laboratories working in the area of aquaculture genetics. Another consideration is the expense of genotyping in relation to sample sizes. Microarray (gene chip) technology and quantitative PCR are particularly useful in medical and clinical settings where large numbers of samples (thousands of individuals per locus) are involved and that can justify the cost involved in the development of the gene chips and hybridization probes. Mass spectroscopy and pyrosequencing are relatively cost-effective (after acquisition of the equipment) when working with relatively small sample sizes (e.g., hundreds of individuals per locus), as is most likely the case with aquaculture and fisheries species.

SNPs can be genotyped with a wide range of techniques and instrumentations, from small-scale, low-budget to expensive high-throughput systems. For SNP genotyping, the greatest determinants of the genotyping platform depend on the availability of equipment. Given the availability of the equipment, considerations can be made based on budget, number of markers, number of individuals, and the requirement for robustness. In spite of its low levels of application in aquaculture and fisheries genome research, SNP markers should gain popularity as more and more sequence information becomes available in aquaculture species. Equally important, once the genetic linkage maps are well constructed, genome scans for QTLs are expected to follow to study traits important to aquaculture, which then depends on the use of well-defined association analysis. As SNP markers are great markers for the analysis of trait-genotype associations, their increased application in aquaculture and fisheries is assured.

Microarray technology

In addition to DNA marker technologies and genome mapping technologies, microarray technology is very important for genome scale analysis of gene expression. This is particularly important for environment-related issues. While microarrays utilize several recent technological innovations, they are, at their core, simply a high density dot blot where DNA samples are applied to a solid support in the form of very small dots, and hybridized to specific DNA probes. Microarrays achieve higher gene feature densities and, therefore, greater power for expression analysis by applying new tools to this old process. High-density spotting robots and photolithography allow each feature to be placed accurately within nanometers of the next feature on a glass slide,
clearly an impossible task with the human hand. Furthermore, fluorescence-based probe labeling provides a cleaner and clearer signal than the radiation traditionally used in blotting. Finally, laser scanners facilitate the resolution of such tremendous feature densities and provide accurate fluorescent signal quantification. Microarray technology allows the changes of gene expression with a specific treatment to be determined at the entire genome scale. For instance, Ju et al., (2002) used microarray technology to determine which genes were up or down regulators in the brain of catfish after treating the fish with cold temperature, and found that 61 genes were significant up-regulators and 12 were down-regulators.

Potential applications of microarray technology in aquaculture and fisheries are wide open. As a genome expression analysis technology, it can be used for analysis of gene expression after any treatment. The first microarray experiment was conducted in catfish for the analysis of cold acclimation (Ju et al., 2002; Kocabas et al., 2004). The microarray created by the GRASP project is widely used in the aquaculture community (Rise et al., 2004a; von Schalburg et al., 2005) for gene expression profiling after infection and vaccination, and stress (Rise et al., 2004b; Purcell et al., 2006; Ewart et al., 2005; Sarropoulou et al., 2005). It is expected that microarrays will find great applications in aquaculture and fisheries.

**Gene mapping technologies**

Although the term gene mapping is widely used in the scientific community, it really refers to several different types of mapping approaches including genetic linkage mapping, physical mapping for the construction of BAC contigs, radiation hybrid mapping, QTL mapping, cytological mapping by FISH (fluorescent in situ hybridization), and comparative mapping. The goal of linkage mapping is to conduct mapping using polymorphic DNA markers in a segregating population (usually F2 population or backcross progenies). Physically linked DNA markers co-segregate. The greater the marker distance, the more likely the recombination during meiosis. Based on co-segregation, markers are placed into the same linkage groups; based on recombinant frequency, marker distances are assigned. Linkage mapping is the basis for genome analysis, and linkage maps have been constructed in many aquaculture and a few fisheries species. The quality of linkage maps are measured by marker density. All aquaculture linkage maps are framework maps or intermediate density maps.

**Quantitative trait loci (QTL) mapping technology**

The goal of QTL mapping is to locate the positions of quantitative trait loci. Most, if not all, performance and production traits of aquaculture are controlled by multiple genes and therefore are inherited as quantitative traits. These genes segregate along with linked DNA markers. By measuring association of trait segregation patterns with marker segregation patterns, it is possible to place trait (or genes responsible for the trait) on linkage maps. QTL studies have been conducted mainly in farmed fish species; for example, rainbow trout. It is expected that QTL mapping will be the key to genetic improvements using marker-assisted selection.

**Physical mapping technology**

Although several approaches are available for physical mapping, the most popular is the BAC-based approach (Bacterial Artificial Chromosome). This approach is based on restriction fingerprinting. Adjacent overlapping DNA segments should share fingerprints that allow large insert BAC clones to be lined up in a linear fashion reflective of their position in the genome.

BAC-based physical mapping has been conducted in several fish species, but mostly with model species. BAC-based physical maps have been only conducted in a few farmed fish; for example, Nile tilapia (*Oreochromis niloticus*) (Katagiri et al., 2005),
Atlantic salmon (*Salmo salar*) (Ng *et al*., 2005), and channel catfish (Xu *et al*., 2007). BAC libraries have been constructed for more farmed species including rainbow trout, Pacific and eastern oysters, and penaeid shrimps. Because physical maps are required for position-based gene cloning, it is expected that physical maps will be constructed soon for many important farmed fish species.

**Other mapping technologies**

Cytological approaches have been used to map genes to chromosomes of some farmed fish species, but because of its relatively low resolution, this mapping strategy is used only as a complementary strategy for the purpose of chromosome marking and related purposes. Radiation hybrid mapping panels have been only established in zebrafish (*Danio rerio*) and European sea bass (*Dicentrarchus labrax*) (Senger *et al*., 2006). Although this approach has been extremely popular in mammalian species, its application in fish is limited. The major reason is that BAC-based physical mapping provides greater levels of resolution and is also more cost effective. The goal of comparative mapping is to use known information from a map-rich species for genome studies of a map-poor species. Knowing the location of genes in a well studied species such as a related model species like zebrafish, one can ask if the genes are arranged similarly in the same chromosomal locations. Comparative mapping is still at its infancy stage in aquaculture species, but hold great promises for the identification of candidate genes responsible for important economic traits.
Fish genetic resources (FiGR) comprise all finfish and aquatic invertebrate genetic material that has actual or potential value for capture fisheries and aquaculture. In capture fisheries more species are becoming endangered and more stocks overexploited. Management of FiGR can help maintain and rebuild these fisheries. Deep-sea fisheries and modern genetic technologies are emerging areas that require attention. Improved information is necessary for improved policies, but at present it is incomplete, scattered and unstandardized. Although tremendous progress has been made in the genetic improvement, genetic stock identification and genomics of aquatic species, further work is needed to: (i) assess the status of FiGR in capture fisheries and aquaculture; (ii) improve the capacities of scientists, technical persons, governments and industry; (iii) improve facilities for characterizing FiGR; (iv) develop genetically improved farmed types of aquatic species; (v) develop appropriate policy instruments on use and conservation of FiGR; (vi) improve general awareness and levels of knowledge about FiGR; and (vii) prioritize species, geographic areas and production systems on which to expend resources for conservation and use of FiGR.