

FAO International Technical Conference

Agricultural biotechnologies in developing countries: Options and opportunities in crops, forestry, livestock, fisheries and agro-industry to face the challenges of food insecurity and climate change (ABDC-10)

Guadalajara, Mexico, 1 – 4 March 2010

Current Status and Options for Biotechnologies in Fisheries and Aquaculture in Developing Countries

Acknowledgements

Preparation of this document was coordinated and finalized by Rohana Subasinghe and Doris Soto of FAO's Aquaculture Management and Conservation Service, with assistance from Preet Lidder and John Ruane of the ABDC-10 Secretariat. Victor Martinez (University of Chile, Santiago) provided the initial draft and John Benzie (University College Cork, Ireland) and CV Mohan (Network of Aquaculture Centres in Asia-Pacific, Bangkok, Thailand) provided valuable technical inputs.

Table of Contents

Acronyms and Abbreviations

1. Fisheries and aquaculture: historical background

1.1 Introduction

1.2 Fish as a protein source

1.3 Aquaculture

A. Stocktaking: Lessons from the Past

2. Overview of main areas where biotechnologies are being applied in aquaculture and fisheries in developing countries

2.1 Genetic improvement and control of reproduction

2.2 Biosecurity and disease control

2.3 Environmental management and bioremediation

2.4 Biodiversity conservation and fisheries management

3. Current status of application of biotechnologies in developing countries

3.1 Genetic improvement and control of reproduction

3.1.1 Polyploidy

3.1.2 Gynogenesis/Androgenesis

3.1.3 Controlling time of reproduction in fish and shellfish

3.1.4 Development of monosex populations

3.1.5 Cryopreservation

3.1.6 Genomics

3.1.7 Genetic modification

3.1.8 Molecular markers

3.2 Biosecurity and disease control

3.2.1 Pathogen screening and disease diagnostics

3.2.2 Vaccines

3.3 Environmental management and bioremediation

3.4 Biodiversity conservation and fisheries management

3.5 Concluding remarks

4. Case studies

4.1 PCR-based pathogen detection in shrimp aquaculture in India

4.2 Specific pathogen-free (SPF) stocks in shrimp aquaculture

B. Looking forward: Preparing for the Future

5. Key issues where biotechnologies could be useful

6. Identifying options for developing countries

7. Identifying priorities for action for the international community

C. References

Acronyms and Abbreviations

AFLPs = amplified fragment length polymorphisms

BMP = better management practice

cGRASP = consortium for Genomics Research on All Salmon Project

ELISA = enzyme-linked immunosorbent assays

ENSO = El Niño-Southern Oscillation

EST = Expressed sequence tag

GMO = genetically modified organism

GnRH = gonadotrophin release hormone

IHHNV = infectious hypodermic and haematopoeitic necrosis virus

IHNV = infectious haematopoeitic necrosis virus

IPNV = infectious pancreatic necrosis virus

ISH = in situ hybridization

MAS – marker-assisted selection

N_e = effective population size

OIE = World Animal Health Organisation

PCR = polymerase chain reaction

PIT = passive integrated transponder

QTL = quantitative trait locus

RAPDs = random amplified polymorphic DNAs

RT-PCR = reverse transcriptase-PCR

SNP = single nucleotide polymorphism

SPF = specific pathogen-free

TSV = Taura syndrome virus

WSSV = white spot syndrome virus

YHV = yellow head virus

1. Fisheries and aquaculture: historical background

1.1 Introduction

Capture fisheries and aquaculture supplied the world with over 113 million tonnes of food fish in 2007, providing an apparent per capita supply of 17.1 kg (live weight equivalent), which is among the highest on record. Global production of fish from aquaculture has grown rapidly during the past four decades, contributing significant quantities to the world's supply of fish for human consumption. Aquaculture currently accounts for nearly half (44.3 percent) of the world's food fish (Figure 1). With its continued growth, it is expected that aquaculture will in the near future produce more fish for direct human consumption than capture fisheries (FAO, 2009).

Started as primarily an Asian freshwater food production system, aquaculture has now spread to all continents, encompassing all aquatic environments and utilizing a range of aquatic species. From an activity that was principally small-scale, non-commercial and family-based, aquaculture now includes large-scale commercial or industrial production of high value species that are traded at the national, regional and international levels. Although production remains predominantly Asian and still largely based on small-scale operations, there is a wide consensus that aquaculture has the potential to meet the growing global demand for nutritious food fish and to contribute to the growth of national economies, while supporting sustainable livelihoods in many communities (Subasinghe, *et. al.* 2009).

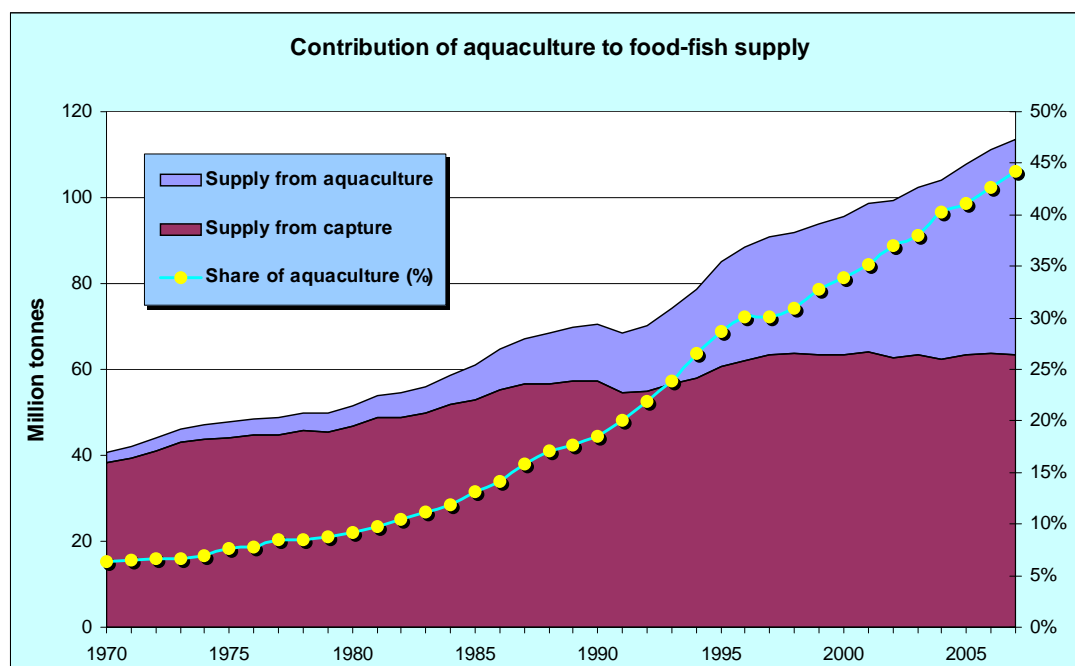
1.2 Fish as a protein source

In 2006, fish provided more than 2.9 billion people with at least 15 percent of their average per capita animal protein intake. The contribution of fish to the total world animal protein supplies grew from 14.9 percent in 1992 to a peak of 16.0 percent in 1996 before declining to about 15.3 percent in 2005. Notwithstanding the relatively low fish consumption in low-income food-deficit countries of 13.8 kg per capita in 2005, the contribution of fish to total animal protein intake was significant – at 18.5 percent – and is probably higher than indicated by official statistics in view of the under-recorded contribution of small-scale and subsistence fisheries and aquaculture (FAO, 2009).

1.3 Aquaculture

Aquaculture is the farming of aquatic organisms, including fish, molluscs, crustaceans and aquatic plants. Farming implies modifications and intervention in the production cycle, such as regular stocking, sorting, feeding and protection from predators in order to enhance production. It is important to note that aquaculture has a long tradition in the developing countries of the Asia-Pacific region, supplying most of the world's aquaculture production (over 90 percent), and making important contributions to the livelihoods and subsistence of small-scale farmers and coastal populations in many countries in the region. In Latin America, small-scale aquaculture has yet to be widely developed; however, there are several examples of newly established industries based on intensive aquaculture practices, especially using exotic species. Salmon farming in Chile is one of the best examples, but there are also expanding aquaculture industries for shrimp and tilapia culture in Ecuador, Costa Rica and Honduras. While Europe and North America import significant quantities of farmed aquatic animals, they also produce fish and shellfish, both from freshwater and marine environments. Africa's contribution to global aquaculture is still small; however, the region is moving forward and increasing production.

Figure 1. Contribution of food fish supply from capture fisheries and aquaculture from 1970 to 2006. (FAO FishStat and FAO, 2009).



Aquaculture covers a wide range of species and methods. It is practised from the cold waters of the far north and south, where fish like salmon, Arctic char and sturgeon are grown in ponds, flowing raceways and cages in the sea, and through the latitudes as far as the tropics, where carp and tilapia flourish in freshwater and shrimp and sea bass are farmed along the coasts. It ranges from the production of fish in naturally occurring ponds in rural areas to the intensive culture of ornamental fish in plastic tanks in the middle of a city. It is practised by the poorest farmers in developing countries as a livelihood and supply of much needed protein for their families, and by urban sports shop owners in Europe and North America producing baitfish for weekend anglers.

1.3.1 Farming systems

Aquaculture systems can range from an intensive indoor system monitored with high-tech equipment through to the simple release of fry and fingerling to the sea, but the aim remains the same: to improve production. Some of the simplest production systems are the small family ponds in tropical countries where carp are reared for domestic consumption. At the other end of the scale are high technology systems, such as the intensive indoor closed units used in North America for the rearing of striped bass or the sea cages used in Chile and Europe for growing salmon and bream.

All products and systems are geared to produce animals for market and are much governed by market demand at all levels. Regardless of whether it is a high-value commodity like shrimp, salmon or grouper, or a low-value commodity such as carp and Tra catfish, all products are destined for markets, be they local, regional or international. All production systems contribute to food security and human development, although small-scale rural production systems provide more support to improving or maintaining livelihoods and generating employment and income for many around the world.

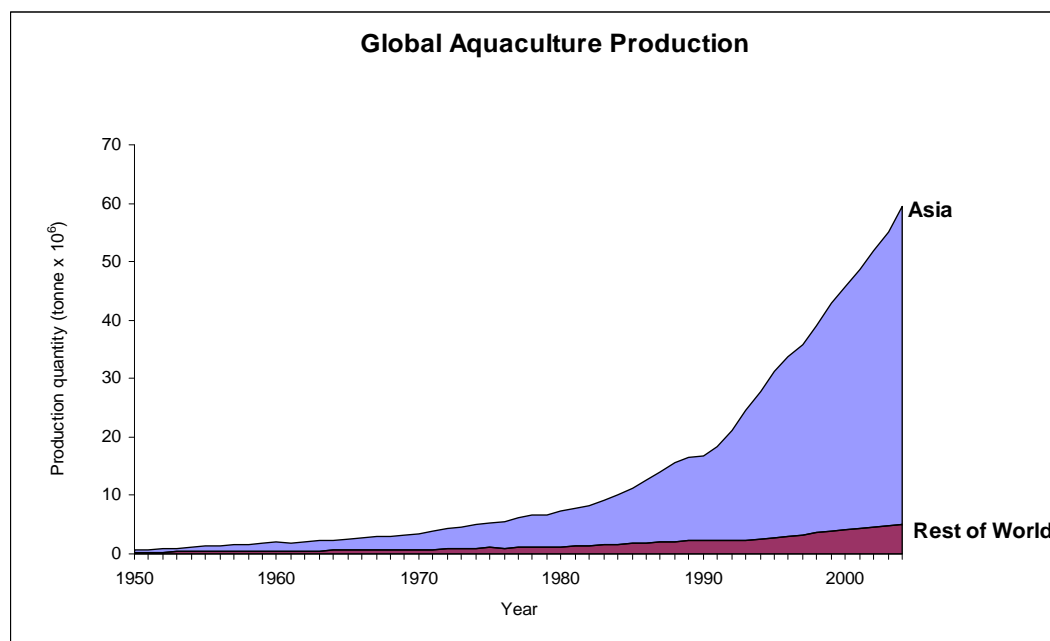
It is important to note that most of these small-scale aquaculture activities occur in developing countries, especially in regions or rural areas where food supply is at risk. For example, tilapia has become a globally important aquatic species that is produced in nearly 100 developing countries worldwide. According to FAO, about 80 percent of the world's farmed tilapia comes from small-holders in developing countries, and this species is particularly prominent in production systems

in the Asia-Pacific, the region that provides most of the world's aquaculture supply (De Silva, Subasinghe, Bartley, 2006).

Another good example of extensive aquaculture is the production of major carps in India. In this case, the majority of the production takes place in rural areas with relatively few impacts on the environment, particularly by using multitrophic culture of species such as catla (*Catla catla*), rohu (*Labeo rohita*) and mrigal (*Cirrhinus mrigala*). It is true that some instances of uncontrolled aquaculture development have caused significant negative environmental and social impacts. However, except for a very few species, there are few negative environmental impacts associated with current production systems and practices. Moreover, most traditional and extensive systems produce fish with little or no negative environmental or social impact.

There has been a steady increase in the growth of aquaculture in developing countries, the rate of growth being twice that of developed nations. The most recent figures for global aquaculture production show that more than 90 percent of total fish production comes from developing countries, particularly China, which contributes about 70 percent of the total global fish and shellfish production (Subasinghe *et al.*, 2009). Aquaculture is thus often one of the most important food production sectors in developing countries, and in many cases it is one of the most important sources of both food and income for rural populations (Figure 2).

Figure 2. Global aquaculture production; Asia vs. rest of the world and China vs. Asia. (FAO FishStat and FAO, 2009).



1.3.2 Extensive vs. intensive aquaculture

Aquaculture practice is an example of a strong continuum of production systems. From a simplest production system with absolutely no inputs and with minimal interventions, aquaculture ranges up to highly sophisticated, fully automated, industrial production systems comprising submerged offshore cages producing large quantities of fish from a single unit. Intensive or extensive aquaculture requires good-quality seed for farming. Seed quality is not only dependent on good hatchery technology, but also on good broodstock with improved genetic quality. The genetic quality of the broodstock and seed used in aquaculture can be improved using biotechnological tools and procedures. There have been some interventions, and good results have been reported.

Modern aquaculture, through the intensification of culture systems and the diversification of both the species cultured and the culture methods employed, often creates an ideal environment for disease-causing organisms (pathogens) to flourish. The expanded and occasionally irresponsible global movement of live aquatic animals has been the cause of transboundary spread of many pathogens, which have sometimes resulted in serious damage to aquatic food productivity. Some of these pathogens have become endemic in culture systems and in the natural aquatic environment, thus making them difficult to eradicate. Since they have become endemic, recurrent pathogen incursions and disease outbreaks occur in farms, making it difficult for the farmers to effectively manage farm health. Instead of implementing effective health management strategies and practices, many farmers opt to use antimicrobials as treatments. There is therefore a need to develop alternate methodologies and tools for maintaining aquatic animal health in aquaculture systems. Such tools and methodologies are generally the result of biotechnological research and several success stories exist. Similarly, biotechnological research has also helped in the improvement of feeds, feeding and nutrition as well as of water quality and the environmental impacts of aquaculture.

The document is divided into two parts: "Stocktaking: lessons from the past" and "Looking forward: preparing for the future". In the "Stocktaking" part, Section 2 provides a brief overview of the main areas where biotechnologies are currently been applied, and Section 3 documents the current status of application of biotechnologies in developing countries. Section 4 presents two relevant case studies. The second part of the document, "Looking Forward", encompasses three sections: Section 5, which looks at a couple of key issues for the future where biotechnologies could be useful; Section 6, which identifies a number of specific options for developing countries to help them make informed decisions regarding adoption of biotechnologies; and Section 7, which proposes a set of Priorities for Action for the international community (FAO, UN organizations, NGOs, donors and development agencies).

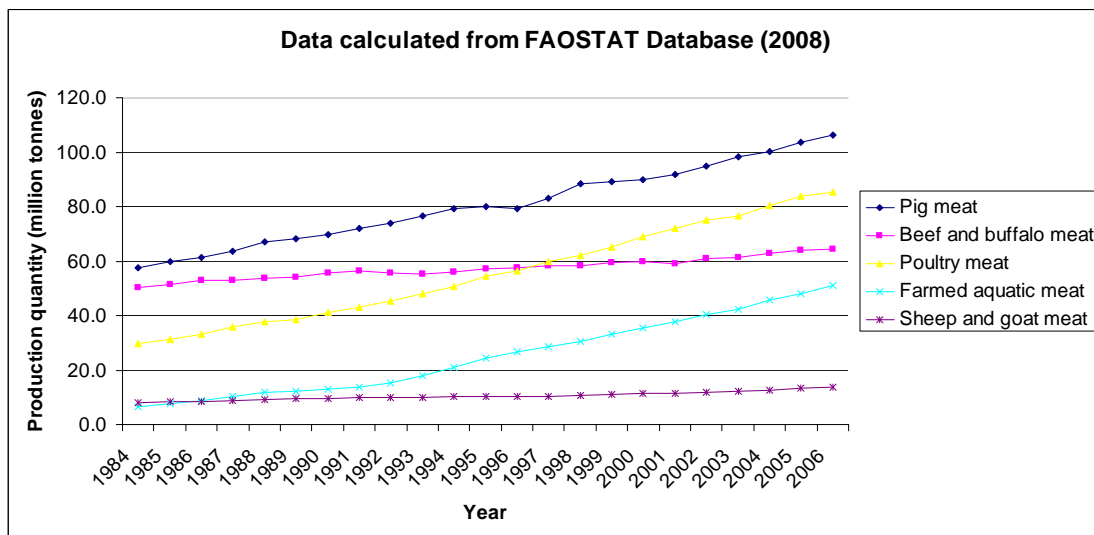
A. Stocktaking: Lessons from the Past

2. Overview of main areas where biotechnologies are being applied in aquaculture and fisheries in developing countries

2.1 Genetic improvement and control of reproduction

Aquaculture is still the fastest growing food producing sector, compared to other food commodities (FAO, 2009) (Figure 3). One of the reasons for this is the diversity of species in culture at present (over 230), and the genetic diversity that can be exploited through captive breeding and domestication, enabling the development of improved culture methods for a diverse array of species to expand commercial aquaculture (Subasinghe, 2009). A lack of knowledge of the biology of many of these species and the cost of technology development are constraints that explain in part why biotechnologies are only now emerging as useful tools for increasing the productivity and sustainability of this sector. Aquaculture is a sector that is likely to benefit greatly from the application of appropriate genetic and reproduction biotechnologies to increase food production.

Figure 3. Growth in production of different food commodities – 1984–2006



Despite the current trend towards the intensification of production systems, aquaculture has not made full use of conventional technologies such as genetic selection and breeding improvement programmes to increase production as other food production sectors have. The rearing of many newly cultured species is to a large extent based on juveniles and/or broodstock obtained from the wild. In order to establish practical breeding programmes to produce seed in hatcheries, it is necessary to have a detailed understanding of the complete production cycle. Such knowledge is also required in order to disseminate breeding improvements to the production sector.

One of the best examples is the inability to fully domesticate *Penaeus monodon*, the black tiger prawn, arguably the most valuable species produced globally. Although specific pathogen-free (SPF) hatchery stocks bred for improved growth have become available recently, production still depends on broodstock collected from the wild. As a result, production of this species has been replaced over the last few years with that from the white shrimp, *L. vannamei*. Improved SPF *L. vannamei* have been readily available for some time, and now supply essentially 100% of farmed white shrimp and more than 60% of all farmed penaeid shrimp world wide. The shrimp

aquaculture sector therefore illustrates the benefits of genetic improvement for increasing production and competitiveness of aquaculture industries.

The *P. monodon* example illustrates how a lack of knowledge concerning some phases of the life cycle, such as reproduction or metamorphosis, may be a limiting factor in developing domesticated stocks. Certain species of tuna, a marine resource that is being harvested under a quota system, are now produced in considerable quantities in captivity or culture. The aquaculture production of this valuable species will undoubtedly increase once the life cycle is closed and the hatchery production of tuna fry becomes a reality. This scenario is also applicable to the hatchery production of mollusc species. There is a huge demand for spat (fertilized shellfish larvae) but most spat are still coming from the wild.

The use of hormones for the control of reproduction has been primarily developed for inducing the final phase of ova production, i.e. for synchronizing ovulation and for enabling broodstock to produce fish in the first part of the season or when environmental conditions suppress the spawning timing of females. These procedures began by the pioneering work of Houssay (1930), who demonstrated that extracts of hypophysis (pituitary gland) can have an effect on sexual maturation of fish and reptiles (Zohar and Mylonas, 2001). These results allowed the development of a relatively simple procedure consisting of injecting hypophysial extracts purified by chromatography that contain products such as inductive hormones related to sexual maturation. Human chorionic gonadotrophin and the gonadotrophin release hormone (GnRH) were also used to control the maturation of many fish species without limitations due to species-specific effects (Zohar and Mylonas, 2001). GnRHa, an analogous GnRH developed chemically, is more efficient in inducing maturation and is relatively inexpensive. It can be injected or administered by means of pellet implants, which facilitates its practical use. The use of hormones such as GnRHa has allowed advancement of the date of egg-laying in several species of fish, mainly salmonids, although for relatively short periods of time (Valdebenito, 2008). Several other molecules are currently under development for use in molluscs (e.g. scallops, oysters and mussels), where synchronous reproduction is required for the hatchery rearing of larvae for aquaculture production in developing countries, instead of using seed obtained from natural banks.

2.2 Biosecurity and disease control

Disease outbreaks are a serious constraint to the development of intensive aquaculture systems and can have a major impact on production due to mortality and decreased growth. It has been recognized that disease is the most significant factor impacting the intensive production of shrimp, salmon, carp and tilapia, with losses of 10 – 90 percent of total production (Peinado-Guevara, Melina and López-Meyer, 2006). Although many aquatic animal pathogens are well studied, unlike in terrestrial animals, the spread of pathogens is easy through water, and control is difficult due to high density culture in fluid environment. Disease occurs in all systems, from extensive to intensive, although heavy losses are always possible in intensive production systems (Bondad-Reantaso, *et al.*, 2005).

Intensive and semi-intensive aquaculture can have important effects on the quality of the aquatic environment in which the animals are reared. Poor water quality resulting from increased waste products, inadequate farm management, increased stocking densities within farms and increased densities of aquaculture units per sector can increase the likelihood of disease outbreaks and other environmental problems such as eutrophication, episodic oxygen shortages, algal blooms, etc., all of them potentially resulting in high mortalities. A more “systems-oriented approach” is therefore needed to provide suitable husbandry for effective growth and to control disease outbreaks effectively.

There is a greater need for management intervention in intensive systems. Here biotechnological tools can be a valuable part of management approaches. Their scope of application is broad – they can be used as sensors in the production environment, for waste management (through controlled microbial technologies), and for disease detection and control (molecular methods). Traditionally, disease control is often carried out only after mortality has been observed. In the past, the

diagnosis of fish diseases has been achieved primarily using histopathological methods, supported by parasitological, bacteriological and viral studies based on necropsy and *in vitro* cell culture. These are well-proven techniques; however, they require a high level of expertise and are often quite time-consuming, not being susceptible to automation. For these reasons, although expert training is required, polymerase chain reaction (PCR) technology (described later) has become an important tool for pathogen assessment in developing countries (in the shrimp industry of Asia and Latin America, for example).

2.3 Environmental management and bioremediation

Aquaculture has often been accused of being unsustainable and not environmentally friendly. Although in some cases, where aquaculture development has failed live up to the global expectations of sustainable development, these allegations are not entirely unfounded, the majority of aquaculture is practised sustainably and with a high degree of environmental conscientiousness. Reducing the impact of effluent discharge, improving of water quality and the responsible use of water are key areas to be considered during aquaculture development. A number of biotechnologies are being used to address these areas: bioremediation for the degradation of hazardous wastes; the use of vaccination and probiotics to reduce antimicrobial use in aquaculture; and the use of DNA-based methodologies for the early detection of toxin-producing algae.

2.4 Biodiversity conservation and fisheries management

In fisheries management, conservation is an important concept. Good fisheries management requires effective conservation measures, which require better understanding of the population structure of the fishery. One of the most important population parameters for assessing the fate of a population is the effective population size (N_e). N_e determines the amount of genetic variation, genetic drift and linkage disequilibrium in populations and can be calculated as half the reciprocal of the rate of inbreeding (Tenesa *et al.*, 2007). There is much concern in fisheries and aquaculture production about the potential loss of genetic variation that may result from the relatively high rates of inbreeding expected in these populations. This is because many fish and shellfish species produce thousands or even millions of fertile eggs from a single female. Due to differences in the biological and environmental factors affecting the survival of individual families, many species show a relatively large variance in family size, further decreasing the N_e (Falconer and McKay, 1996). Fisheries resource managers have focused on the actual number of individuals in a population (census numbers) (Grant, 2007), which may be many times higher than the effective population size (Hauser *et al.*, 2002; Primmer, 2006). Therefore, it is difficult or even impossible in some cases to infer the effective population size using the census number. Inadequate procedures for stock enhancement can yield a very small effective population size due to the high prolificacy of fish and shellfish species. Thus a very small number of breeders could be used for restocking purposes, and bottlenecks can affect the fitness of the population in future generations. A range of biotechnology-based approaches are being used to conserve wild fish populations, such as the use of molecular markers: to estimate N_e in wild populations; to study gene flow between farmed and wild fish populations; and to monitor and understand changes in wild fish population sizes (Hansen, 2008; Primmer, 2006).

3. Current status of application of biotechnologies in developing countries

In fisheries and aquaculture, although perhaps not as much as in livestock production and crops, some biotechnologies have been used in developing countries. As mentioned earlier, use of biotechnologies in fisheries is very limited whilst in aquaculture biotechnologies are represented in a few fields such as genetic improvement, disease control, feeds and nutrition and environmental improvement.

3.1 Genetic improvement and control of reproduction

3.1.1 Polyploidy

Many fish and shellfish species are relatively tolerant to chromosomal manipulation in the early stages of their development. The use of genetic manipulation, including polyploidy (i.e. increasing the number of sets of chromosomes), to improve aquaculture production has been examined. However, there has been little discussion of the use of these technologies in practical management programmes in developing countries, or on how they can be used efficiently within the context of breeding programmes. Furthermore, the potential value of this technology under practical conditions for enhancing the performance of commercial populations in developing countries is not clear.

The induction of polyploidy has been considered by many researchers (Purdom, 1983; Thorgaard, 1986) because of the advantages related to triploid sterility. For example, triploids (with three sets of chromosomes) may be useful for conservation programmes, where sterility can prevent introgression of genes from escaped individuals from commercial stocks into natural populations (Galbreath *et al.*, 1994) or in commercial operations, where sterile fish are desirable to prevent the side effects such as deterioration of carcass quality due to maturation or the occurrence of high mortalities in stocks when males mature early or that occur prior to maturation, especially in populations Pacific salmon (Purdom, 1983; McGeachy *et al.*, 1995).

Triploidy leads to the production of nearly completely sterile populations, as has been observed in rainbow trout populations with spontaneously occurring triploids (Thorgaard and Gall, 1979). However, the degree of reproductive disruption varies depending on the species and the sex. Gametogenesis is severely disrupted in triploid females of salmon, while in contrast, triploid males usually display secondary sexual dimorphism (i.e. darkened skin colour and modified body conformation), courtship behaviour, and develop an endocrine profile similar to that of diploid males. Spermatogenesis, however, appears to be somewhat reduced in comparison to diploid males (Benfey *et al.*, 1986). Although triploid males are to a great extent sterile, fertilization has been reported to occur. In the salmon aquaculture industry, sexual maturity and the associated gonadal development is generally an economic drawback, as metabolic energy is diverted from somatic cell growth to reproduction, resulting in the deterioration of flesh quality and appearance. In this situation, the advantages of triploidy occur primarily after the onset of maturation, when triploid female fish may show an extension of growth (Thorgaard, 1986) and the inhibition of maturation prevents the normal degradation in carcass quality that is observed during the spawning season (Asknes *et al.*, 1986). Furthermore, female salmon triploids show a significantly higher dress-out percentage (Thorgaard and Gall, 1979) and higher pigment (canthaxanthin) retention (Choubert and Blanc, 1989), but concomitantly, there is an increase of fat deposition surrounding the viscera.

In developing countries, the practical implementation of triploidy in fish production has not been very successful. Most of the research on the application of this biotechnology has been experimental, without extensive testing under practical conditions that consider the wide range of environments in which aquaculture takes place. In species such as tilapia and carp, testing of triploidy is a very important issue considering that there is intraspecies variation in the rate of triploidization due to the size and quality of the eggs. For this reason, it is not possible to ensure 100 percent triploidy when applying this technique on a commercial scale. Also, an increased mortality rate at the beginning of the life cycle and the detrimental effect of triploidy on growth

and fitness could be a significant constraint to the commercial production of triploids in some species (Basant *et al.*, 2004). The lack of knowledge about the effects of competition between triploids and diploids in large extensive conditions in species such as tilapia could also be a disadvantage, since triploids sometimes lack robustness compared to normal diploids, but this expression varies among species (Benfey, 1999). In many cases, the variation in performance between diploid and triploid stocks has not been fully estimated, and thus it may not be possible to accurately predict the relative performance of triploids in commercial conditions, which may be a problem in conventional breeding programmes of many fish and shellfish species (Pechsiri, 2007).

In developing countries, for various reasons these techniques are not currently used for commercial purposes. Tilapia, for example, cannot be easily reproduced using external fertilization, which is a prerequisite for shock treatment. Furthermore, when a very small number of eggs is obtained per spawn, it is not possible to ensure a constant rate of triploidy per spawning. In rainbow trout, it is only profitable to use triploid females, since males show some degree of reproductive onset. For developing such female triploid populations, neomales (i.e. morphologically female but genetically male, see below) are required, which in some instances are difficult to stock up to a commercial scale. In Indian carps, sterility aiming at faster growth and thus enhanced production may not be cost-effective, since harvesting after one year of age is not profitable (males mature at one year of age and females when approaching two years).

In southern India, precocious maturation is a potential constraint on yields of cultured common carp, as both males and females can attain sexual maturity well before reaching a marketable size. However, triploid fish did not show any improvement over diploid individuals, except for higher dress-out percentages (Basavaraju *et al.*, 2002).

In spite of the fact that there has been a plethora of research conducted on triploidization and chromosomal biotechnologies, there remains a gap between research findings and the practical implementation of triploidy. Several reasons explain this fact. The usefulness of applying chromosomal biotechnologies such as triploidy for aquaculture production seems to be very species specific, and therefore in some cases (such as in salmon, tilapia and carp), the advantages due to delayed maturation or increased growth are unclear. Furthermore, the results of using these techniques to increase growth rate or delay reproduction are not seen as sufficiently beneficial for the technique to be implemented on a large scale (P. Routray, Central Institute for Freshwater Aquaculture, personal communication, 2009).

For the technology to be practical, it should be possible to produce all-triploid populations without the need to test the triploidy status of each batch of embryos produced. Because triploidy induction using thermal shock is not 100 percent effective, this is a serious drawback to the large-scale commercial application of the technique. Crossing between tetraploids and diploids is a way to produce 100 percent triploids; however, in most species tetraploid production is not straightforward. Furthermore, the genetic lag between the tetraploid population and the diploid breeding programme can seriously affect the efficiency of the production system. For all these reasons, this technology has not been used extensively in developing countries for production purposes.

3.1.2 Gynogenesis/Androgenesis

Gynogenesis is the production of an embryo from an egg after penetration by a spermatozoon that does not contribute genetic material. Androgenesis is the production of an embryo from an egg whose DNA was inactivated and which was fertilized using normal sperm. In both cases, the diploidy is restored by using heat/cold shocks. In gynogenesis, if diploidy is restored soon after fertilization, the procedure is called meiotic gynogenesis, due to the fact that the second polar body is retained, and this procedure is similar to what is expected under autofertilization in terms of inbreeding. If shocks are applied later or in androgenesis, where the ova was DNA-irradiated for DNA inactivation, the same chromosome is duplicated, and thus the embryo is a double haploid individual, which is completely inbred for every loci.

Several papers have discussed the usefulness of this type of reproduction for genetic analysis in carp, tilapia and rainbow trout breeding programmes. In some cases, the use of gynogenetic individuals has been suggested for capitalizing on non-additive genetic effects, to increase additive genetic variance and for product uniformity (Bijma *et al.*, 1997). However, the production of gynogenetic lines is not without problems; after a first round of gynogenesis from an out-bred population deleterious and/or lethal effects can be fully expressed in the double haploid progeny, which may be a problem when implementing a breeding programme from this source. Furthermore, phenotypes cannot actually be a direct reflection of the same trait measured on normal progeny due to developmental instability. Therefore, the utility of this type of reproduction for practical use in breeding programmes is seen as risky in most cases. Nonetheless, they can be used effectively for developing powerful quantitative trait locus (QTL) mapping experiments using the surviving clonal lines of this sort obtained from an out-bred population, but this requires having available the gynogenetic lines that are needed for further assessment (Martinez, 2007).

3.1.3 Controlling time of reproduction in fish and shellfish

So far the application of hormonal treatment has been quite successful, especially for controlling reproduction in broodstock. This is specially the case in salmon and trout farming in Chile, where either implants or injection of the hormonal compound are used extensively in salmon farming for synchronizing reproduction. Since hormone application is not done in the commercial fish, but rather in the broodstock, which are discarded for human consumption, these procedures are not subjected to a negative consumer preference. In carp breeding, the use of hormones has made it possible to artificially manipulate the number of times and the timing of spawning of major Indian carps and African catfish (Routray *et al.*, 2007).

3.1.4 Development of monosex populations

One of the major constraints in practical programmes in developing countries is the fact that mixed sexed populations can behave poorly in production conditions (Subasinghe *et al.*, 2003). This is primarily due to the negative side-effects of early reproductive onset that decrease the growth rate through a series of physiological mechanisms. The faster growth rate of the other sex is probably caused by its later maturation. The negative relationship between growth rate and gonadal development has been found in many species. One explanation of this finding is the appearance and accumulation of sex hormones that act as a growth inhibitory agent (Hulata *et al.*, 1985).

The advantages of monosex culture depend on the species involved (FAO, 1995). This is because one sex may be superior in growth or have a more desirable meat quality, or to prevent reproduction during grow-out or the appearance of sexual/territorial behaviour (aggressiveness) that occurs when a mixed sexed group triggers the reproductive season. For example, female sturgeon are more valuable than males because they produce caviar; female salmon are more valuable because sexually precocious males die before they can be harvested, and salmon roe has an economic value; and male tilapia are more desirable than females because they grow twice as fast and because reproduction is not significant in males during grow-out.

The sex of fish can easily be manipulated using hormonal treatments. In many fish and shellfish species, sex is not permanently defined genetically and can be altered by a number of factors, including hormonal treatment during the early stages of development. Gonadal development starts from primordial germ cells, with females starting differentiation prior to males (Phelps, 2001). The point in time when differentiation occurs depends on the species involved. In tilapia and trout, this mechanism is triggered early in life, while in grass carp and paddlefish it is the opposite (Phelps, 2001). Considering this pattern of development, treatment with the steroid methyl testosterone can be used to develop all-male tilapia populations (Mair, 1999) and androgens (male sex hormones) can be used in trout and carp monosex culture.

There has been concern about the use of hormones in animal production, including in aquaculture systems, that stems from the risk of presence of residues in final products. In spite of the fact that there is little evidence regarding hormonal residues in fish whose sex has been reversed early in life, consumer acceptance may be compromised as a result of the perception of hormonal treatment itself (Subasinghe *et al.*, 2003). For this reason, it appears that other biotechnologies have had more use in developing countries whose production goes mainly to export markets. A variation on this scheme is to produce all male progeny in one more generation. This requires feeding young fish with estrogens (female sex hormones), resulting in a population of all female fish (Fitzsimmons, 2001). These morphologically female but genetically male fish (neofemales) are then raised to maturity, when they are mated to normal male fish. After maturation, the all-male fry produced are tested in order to identify the “super males” (YY), which are then crossed to normal females (XX), thus generating all true male (XY) progeny. The importance of this method is that male fry for commercial production can be produced that have never been treated with hormones. However, one of the disadvantages is that this technique requires more than a single generation to obtain the all-male fry, i.e. this procedure cannot be used without extensive progeny testing to determine which “female” fish will produce all-male progeny, thus requiring a reasonable time span for developing the neomales.

Although tilapia breeding programmes using YY super males are possible, this procedure is not necessarily required, because the application of direct hormonal treatment of undifferentiated fry to produce monosex populations is still a major breakthrough. However, the great expansion of tilapia aquaculture in Asia has been due to mixed-sex tilapia culture, which addresses the high demand for relatively small fish (i.e. fish less than 300 g) that can be obtained by rearing the highly selected genetically improved farmed tilapia and other strains.

3.1.5 Cryopreservation

The aim of the cryopreservation of gametes is related to:

- Disseminating semen from males obtained from selection programmes showing significant response;
- “Refreshing” commercial populations in order to avoid the negative impact of bottlenecks;
- Directly assessing the rates of genetic gain in ongoing breeding programmes; and
- Making semen available across the reproduction window when asynchrony of reproduction exists between males and females (usually males mature earlier than females).

Sperm cryopreservation has been successfully implemented for a number of cultured finfish and shellfish species, and modest success has been achieved in the cryopreservation of shellfish embryos and early larvae. Cryopreservation of finfish ova and embryos has not been successful, which is a major difference with respect to terrestrial animals. This is mainly due to the size of the ova, which are usually large and have thick chorionic membranes that do not facilitate the inclusion of cryoprotectors.

The use of cryopreserved gametes for commercial purposes is still very limited in developing countries. One explanation for this finding is that this biotechnology may require specialized labour and automated procedures to decrease variability in success rates among batches of sperm. Furthermore, it is still uncertain whether this method is economically advantageous compared to disseminating improved broodstock using larval material. In spite of this, the technology has been used for disseminating improved “Jayanti” rohu in India and for the dissemination of improved semen in Sri Lanka (Routray, personal communication). In rainbow trout, cryopreservation has been used for storing semen from neomales, but the problem of highly variable fertilization success remains.

3.1.6 Genomics

Genomics is the study of the genomes of organisms. It includes the intensive efforts to determine the entire DNA sequence of organisms via fine-scale genetic mapping.

Genome Sequencing

One of the major constraints in the rearing of many different aquaculture species is the lack of adequate genomic information. This is due to the fact that sequencing all the species currently used in aquaculture would be costly. Productive species currently being sequenced are the tilapia and the Atlantic salmon (*Salmo salar*). A multinational initiative for Atlantic salmon aims to sequence the genome using Sanger sequencing in order to obtain a genome coverage of more than 6 fold. The project is a partnership between Canada, Norway and Chile, countries that are interested in applying this sequence data for studies related to conservation and production enhancement. The project's output will be delivered to the public domain and provide the required genomic resources for developing single nucleotide polymorphism (SNP) chips that will help implement marker-assisted selection programmes in Chilean salmon aquaculture.

Functional genomics

The recent availability of massive amounts of information from functional genomics, such as microarrays, used to assess gene expression or sequence polymorphisms, has contributed significantly to the genomic biotechnology in aquaculture. Two colour microarrays have been developed for salmonid species that are publicly available and are currently being used to assess disease resistance traits in salmon and for candidate gene discovery. In shrimp, several platforms have been devised in China, Australia, Taiwan POC, Singapore and also the United States of America (Wilson and de la Vega, 2005).

The main use of this resource has been to study differential expression of the transcriptome after viral or bacterial acute infection, but also as bioindicators for assessing chronic disease response. Microarrays are being applied to the fields of ecotoxicology and nutrigenomics. For example, gene expression analysis has been used for assessing the effect of pre-challenging white spot syndrome virus (WSSV) on different genes in order to investigate the immunological mechanisms behind the genetic resistance and to assess potential genes explaining disease resistance at the experimental level in the culture of Pacific whiteleg shrimp (*Litopenaeus vannamei*) in Colombia. In Chile, the salmon microarray available for the consortium for genomics research on all salmon project (cGRASP, <http://web.uvic.ca/grasp/>) in Canada has been used in collaboration with the University of Victoria for assessing disease resistance of piscirickettsia and infectious pancreatic necrosis virus (IPNV) in Atlantic salmon.

3.1.7 Genetic modification

A genetically modified organism (GMO) is one whose genetic material has been altered through genetic engineering techniques with DNA molecules from different sources that are combined into one molecule to create a new set of genes. Typically, it involves introduction of a single gene from an unrelated species. After about two decades of very intensive research, the technology has reached the stage where it is possible to produce GM carp, tilapia and salmon. However, no aquatic GMOs have yet been approved for commercial release for food and agriculture purposes in any developed or developing country. There are potential concerns about the environmental impact of raising such fish (e.g. effects of possible interbreeding with native populations) and the greater amount of feed required for sustaining the increased growth rates, as well as problems with consumer acceptance, which may be one of the most important reasons that transgenic technology has not developed beyond the experimental phase. Many developing countries have yet to develop a clear policy on the use of transgenic fish.

3.1.8 Molecular markers

Marker systems

Molecular markers are identifiable DNA sequences, found at specific locations of the genome, transmitted by standard Mendelian laws of inheritance from one generation to the next. They rely on a DNA assay and a range of different kinds of molecular marker systems exist, such as restriction fragment length polymorphisms, random amplified polymorphic DNAs (RAPDs), amplified fragment length polymorphisms (AFLPs) and microsatellites. The technology has

improved in the past decade and faster, cheaper systems like single nucleotide polymorphisms (SNPs) are increasingly being used. The different marker systems may vary in aspects such as their technical requirements, the amount of time, money and labour needed and the number of genetic markers that can be detected throughout the genome (reviewed in detail in FAO, 2007). RAPDs and AFLPs have been used extensively in aquaculture due to their relatively easy development, i.e. they do not require construction of genomic libraries for their development. Microsatellite markers are used increasingly in aquaculture species (see the review by Liu and Cordes, 2004), due to their elevated polymorphic information content, codominant mode of expression, Mendelian inheritance, abundance, and broad distribution throughout the genome (Wright and Bentzen, 1994).

Molecular markers are being applied in developing countries in both aquaculture and fisheries management. Here, an overview is provided on their use for parentage analysis and genetic selection in aquaculture and for fisheries management and stock enhancement.

Parentage analysis

Molecular markers can be used successfully to trace alleles inherited by progeny from a group of candidate parents, thus providing a means of parentage analysis. In many fish and shellfish species, reproduction cannot be fully controlled and thus natural mating is the only way to produce offspring for the next generation of a breeding programme. For example, tilapia and carp breeding typically involves mass spawning where males and females are stocked in large “hapas” suspended in ponds, where a relatively large number of parents are spawning simultaneously. Since constrained rates of inbreeding are required for sustained rates of genetic gain, in uncontrolled mating schemes it is not always possible to control the genetic contributions of broodstock nor, therefore, the rates of inbreeding in a breeding programme using pedigree information. Small sample sizes, together with sperm competition (Withler and Beacham, 1994), mating preference (as in *Artemia*) and other biological factors after fertilization can increase the variance of family size, thereby decreasing the effective population size to unsustainable levels (Brown, Woolliams and McAndrew, 2005).

When it is possible to control matings, one of the most important constraints still facing effective breeding programmes of species such as salmon, carp and trout is that newborn individuals are too small to be tagged individually using the traditional marking systems for livestock. The application of sustainable breeding programmes requires tagging a constant number of individuals from each family with passive integrated transponders (PIT tags) when they become sufficiently large after a period of individual family rearing, in order to manage the rates of inbreeding. However, this system of early management creates common environmental effects for full-sib families (Martinez *et al.*, 1999). To address these issues, mixtures of equal-aged progeny from different families can be reared communally to preclude the development of such family-specific environmental effects, and genetic markers can be used subsequently to assign individuals to families after evaluation of individual performance (Doyle and Herbinger, 1994). Thus, the impact of early common environmental effects is considerably reduced if markers are used for parentage analysis when selecting individuals for early growth rate traits (Herbinger *et al.*, 1999; Norris, Bradley and Cunningham, 2000). Several multinational salmon companies are using this system of tagging, but there is still no information regarding its economic value compared to conventional tagging systems such as PIT tags. This may be important in species such as carp and tilapia, where the costs of genotyping can greatly outperform the use of tanks and individual tagging systems. Furthermore, it is expected that rates of genetic gain for economic traits will not be significantly affected when common environmental effects are present.

Even though there is a plethora of information in the scientific literature on the use of markers for parentage analysis in fish and shellfish, this procedure has not been fully used in species such as tilapia in developing countries, where basic conventional breeding programmes have proved very successful (Ponzoni *et al.*, 2006). The sample size (i.e. the numbers of individuals and markers required for accurately reconstructing the pedigree of a population) is a practical issue, since not all individuals in a population can be genotyped for all markers available. Issue of sample size

may also arise in species where physical tagging is not possible or not economically sound (e.g. shrimp or marine species) or when disease challenges (e.g. with infectious pancreatic necrosis) are carried out very early stages in the life cycle.

For most breeding programmes, physical tagging will prove efficient both in economic and biological terms to achieve acceptable rates of genetic gain, while minimizing rates of inbreeding. Genetic marker technology can still be costly in developing countries for routine assignment of parentage, although these costs can be reduced using multiplex PCR technology in which more than one marker can be genotyped simultaneously in a single gel lane or capillary (Paterson, Piartney and Knox, 2004). This is especially the case when only DNA markers are used without physical tagging, since individuals must be re-typed when records for multiple traits are included in the selection criteria (Gjerde, Villanueva and Bentsen, 2002). When it is possible to isolate families, multistage selection offers the possibility of first selecting individuals on a within-family basis directly from tanks or hapas (for traits influenced by common environmental effects) and then selecting at a second stage for traits measured at harvest. This alternative would maintain the rates of gain while decreasing the costs associated with tagging, or even increase rates of gain, when recording traits such as body weight from tanks (within families) that can be carried out relatively inexpensively (Martinez *et al.*, 2006).

Marker-assisted selection

Molecular markers can also be used in genetic improvement through so-called marker-assisted selection (MAS), where markers physically located beside (or, even, within) genes of interest (such as those affecting growth rates in salmon) are used to select favourable variants of the genes (FAO, 2007). MAS is made possible by the development of molecular marker maps, where many markers of known location are interspersed at relatively short intervals throughout the genome, and the subsequent testing for statistical associations between marker variants and the traits of interest. In this way, genes (called QTLs) thought to control quantitative traits (traits of agronomic importance controlled by many genes and many on-genetic factors, such as growth rate in fish) can be detected.

MAS can enhance rates of genetic gain compared to conventional breeding for traits that are difficult or expensive to measure or when the heritability is relatively low. So far, many QTLs have been identified in different experiments involving trout, salmon, carp and tilapia, but the main problem of the actual use in MAS is to have enough replications or powerful experiments in order to validate that the actual QTLs detected in a given experiment are real, and are segregating across populations or crosses. Furthermore, many of the QTLs detected were discovered using dominant markers such as RAPDs, which are very difficult to replicate in different laboratories, basically due to the use of insufficient sample sizes and failure to account for the presence of false positives. This outcome is explained by the fact that there is a lack of complete genome sequences for many of the species currently used in aquaculture in developing countries, such as tilapia, carp and shrimp. This is an important practical issue, because, without information from physical maps, it may be difficult to characterize the actual genes explaining the genetic variation explained by the QTLs. This situation reflects the relatively high level of financial resources needed both to carry out a genome sequence project for many species used in aquaculture and to actually implement a MAS programme. This is a very important issue in developing countries where smallholders are less likely to have the financial revenue to allow breeding programmes that incorporate the use of molecular information. Although MAS is potentially useful for many cultured species, conventional breeding programmes may be more profitable in the short to medium term in developing countries in low-input environments.

The development of molecular markers and linkage maps can greatly help scientists to understand the different factors that influence the expression of quantitative traits. A number of genetic linkage maps have been published in aquaculture, some of the most comprehensive being for rainbow trout (Young *et al.*, 1998; Sakamoto *et al.*, 2000; Nichols *et al.*, 2003), channel catfish (Waldbieser *et al.*, 2001), tilapias (Kocher *et al.*, 1998; Lee *et al.*, 2005), Japanese flounder (Coimbra *et al.*, 2003) and mussels (Lallias *et al.*, 2007). In shrimp, recent mapping has

demonstrated the nature of sex control in shrimp as WZ/ZZ like chickens and unknown until now. Still, in important species such as Indian Major carps and Chinese carps they have not been developed. There are a number of ways in which this information can be used, the difference between them being the level of resolution with which these factors can be mapped. For example, QTLs with major effects on quantitative traits are mapped using markers to track the inheritance of chromosomal regions in families or in inbred line crosses using the extent of linkage disequilibrium generated in the population.

In practice, the identification of genes influencing specific traits is achieved using a combination of genetic mapping (linkage and fine mapping) to localize the QTL to a small region on the chromosome under analysis, and candidate gene or positional cloning approaches are used to identify the genes within the QTL region. According to the literature survey, it appears that very little information has come from developing countries on such research issues.

In some cases, it is possible to use sufficient biochemical or physiological information to investigate the association between the quantitative genetic variation and the level of marker polymorphisms within specific genes. Nevertheless, this approach requires a great amount of detailed information in order to choose which gene explains the greatest effect and to have sufficient power to detect the association. This information is starting to appear in the aquaculture literature from multinational projects such as cGRASP, but it is still scarce for other fish species of interest in developing countries.

So far, QTL mapping in aquaculture using commercial populations has been carried out mainly in developed countries, mostly with single-marker analysis (microsatellites and AFLP markers) and using relatively sparse linkage maps when interval mapping is used. In tilapia, the F2 design and a four-way cross between different species of *Oreochromis* have been used for detecting QTLs affecting cold tolerance and body weight (Cnaani *et al.*, 2003). In outbred populations of salmonids, QTLs that influence body weight have been mapped (Reid *et al.*, 2005).

Studies seeking linkage of markers to traits amenable to MAS, such as disease resistance, have begun to appear in the literature over the past few years. For example, QTLs for resistance have been mapped for IPNV in salmonids (Ozaki *et al.*, 2001; Moen, 2007; Houston *et al.*, 2008), infectious salmonid anaemia, infectious haematopoietic necrosis (Rodriguez *et al.*, 2004; Khoo *et al.*, 2004) and stress and immune response (Cnaani *et al.*, 2004) and cold tolerance in tilapia (Moen *et al.*, 2003). Also, Somorjai *et al.*, (2003) reported evidence of QTLs for upper thermal tolerance in salmonids, with differing effects in different species and genetic backgrounds. To date, there are no examples of the application of these QTLs in practical fish and shellfish breeding programmes in developed or developing countries.

3.2 Biosecurity and disease control

Like other farming systems, the aquaculture industry has been overwhelmed by a fair share of transboundary aquatic animal diseases caused by viruses, bacteria, fungi, parasites and other undiagnosed and emerging pathogens. Disease has thus become a primary constraint to the culture of many aquatic species, impeding both economic and social development in many countries. As a result, there will be increasing demand for improved aquatic animal biosecurity, particularly addressing the emerging health problems, based on risk analysis. Epidemiological studies generate the data required for risk analysis; biosecurity measures require good information for accurate assessment, and this leads to appropriate risk management. Thus, biosecurity, risk analysis and epidemiology are highly interrelated. All are aimed at making good use of scientific research for disease prevention, control and management.

Of equal importance is the need for fundamental information that characterizes diseases in aquaculture. Import risk assessment will of necessity set the risk as “high” when there are little data on modes of transmission, host susceptibility, tolerance to abiotic factors (e.g. temperature, salinity) and immune response elicited, for a particular pathogen under consideration. The clear, unambiguous and rapid detection and identification of potential pathogens, using morphological

and molecular diagnostic tools, is of paramount importance prior to making decisions on the disease status of any aquaculture zone.

Although conventional disease-control strategies focus largely on disease control through diagnosis and therapy, the prevention of disease through vaccination, immunostimulation, the use of probiotics and bioremediation in culture environments, nutritional improvements, etc., has also been practised. Significant advancements in these areas have been achieved using biotechnological approaches.

Given the taxonomic diversity of aquaculture species there is also a need to develop better information on the response of these species to disease to develop management strategies for them. Here biotechnology approaches are sometimes the only means by which tools for this can be developed.

3.2.1 Pathogen screening and disease diagnostics

The control of disease outbreaks relies heavily on having rapid and accurate diagnostic tools available in order to detect and identify the pathogen causing mortality. DNA and RNA methods have been used extensively for detecting a number of viral and bacterial pathogens in aquaculture worldwide. The techniques rely upon the fact that each pathogen species carries a unique DNA or RNA sequence that can be used for identification. The techniques offer high sensitivity and specificity, and the commercial development of PCR primers and diagnostic kits allows rapid screening for a number of serious viral and bacterial infections and has direct application. Molecular-based techniques such as PCR also have an application in situations where the animal shows no antibody response after infection. For example, as molluscs do not produce antibodies, antibody-based diagnostic tests have limited application to pathogen detection in these species.

Considering the difficulties that developing countries may face in using advanced molecular diagnostics, and the importance of gradually improving national diagnostic capacities in developing countries, FAO recommended a three-level diagnostic process (FAO/NACA, 2000). This involves: field observations and necropsy (Level I); laboratory observations, bacteriology and histopathology (Level II); electron microscopy, molecular biology and immunology (Level III). In countries where Level II and Level III diagnostic capabilities are not found, initial disease screening is carried out using Level I gross clinical examination. Accompanied by histopathology, this has been the traditional method of detecting pathogens in both developed and developing countries. There is a clear need to improve national diagnostic capacities to reach Level II and Level III diagnosis procedures, including molecular diagnostics.

These tools include both immunoassay and DNA-based diagnostic methods, e.g. fluorescent antibody tests, enzyme-linked immunosorbent assays (ELISA), radio-immunoassay, in situ hybridization (ISH), dot blot hybridization and PCR amplification techniques. They are currently used to screen and/or confirm the diagnosis of many significant pathogens of cultured finfish, such as channel catfish virus, infectious haematopoietic necrosis virus, IPNV, viral haemorrhagic septicaemia virus, viral nervous necrosis virus and bacterial kidney disease, as well as shrimp diseases such as WSSV, yellow head virus (YHV), infectious hypodermic and haematopoietic necrosis virus (IHHNV) and Taura syndrome virus (TSV) (Walker and Subasinghe, 2000). Similar tools are under development for molluscan pathogens (*Haplosporidium* sp., *Bonamia ostreae*, *Marteilia refringens* and Herpes virus). Immunoassays and nucleic-acid assays provide quick results, with high sensitivity and specificity at relatively low cost, and are particularly valuable for infections that are difficult to detect (e.g. subclinical infections) using standard histology and tissue-culture procedures. Molecular tools are also useful for research into the pathology and immunology of specific infections. They can be used with non-lethal sampling and are valuable to monitor challenge experiments under controlled laboratory conditions. Further development of this technology is likely to speed up the detection (field monitoring and laboratory examination) and diagnosis of disease, which is crucial for early and effective control of emergent disease situations.

Antibody-based techniques

A variety of antibody-based tests and molecular tests have been developed to detect mainly bacterial and viral fish pathogens, although tests have also recently been reported for parasites and fungal agents. The antibody-based tests include slide agglutination, co-agglutination/latex agglutination, immunodiffusion, direct and indirect fluorescent antibody tests, immunohistochemistry and ELISA, dot blot/dip stick and western blot. The antibody-based test selected for the identification of pathogens depends on a variety of factors, since each method has its merits and disadvantages. Although such methods are useful for the detection of pathogens in pure culture or/and in infected fish tissue, their sensitivity thresholds limit use in environmental samples, especially where pathogen levels are extremely low. DNA detection methods, however, such as PCR and ISH are ideally suited.

DNA-based techniques

Molecular technologies are also widely used for the detection of fish pathogens (Adams and Thompson, 2008; Adams and Thompson, 2006). They have been successfully utilized for the detection and identification of low levels of aquatic pathogens. Such methods are also particularly useful for micro-organisms that are difficult to culture, may exist in a dormant state, are involved in zoonosis, or in the elucidation of pathogen life cycles. In addition, molecular methods can be used for the identification of pathogens to species level (Puttinaowarat *et al.*, 2000) and in epidemiology for the identification of individual strains and differentiating closely related strains (Cowley *et al.*, 1999). Because of the general unavailability of the traditional pathogen isolation methods and immunodiagnostics for molluscs and crustaceans, molecular techniques have increasingly been used (Berthe *et al.*, 1999; Lightner, 1996; Lightner and Redman, 1998). The DNA-based methods such as PCR are extremely sensitive. However, false positive and false negative results can cause problems due to contamination or inhibition (Morris *et al.*, 2002). Real-time PCR (closed tube to reduce contamination) and nucleic acid sequence based amplification are alternatives that reduce this risk and offer high sample throughput (Overturf *et al.*, 2001; Starkey, *et al.*, 2004). Some of the most common PCR-based technologies used for the detection of pathogens are nested PCR, RAPDs, reverse transcriptase-PCR (RT-PCR), reverse cross blot PCR and RT-PCR enzyme hybridization assay (Cunningham, 2004; Puttinaowarat *et al.*, 2000; Wilson and Carson, 2003). ISH is also widely used in the detection of shrimp viruses (Lightner, 1996; Lightner and Redman, 1998; Tang and Lightner, 1999; Tang *et al.*, 2005) and in the confirmation of mollusc parasites (Stokes and Bureson, 1995; Le Roux *et al.*, 1999; Cochenec *et al.*, 2000; Carnegie *et al.*, 2003). Colony hybridization has also been used successfully for the rapid identification of *Vibrio anguillarum* in fish (Aoki *et al.*, 1989) and has the advantage of detecting both pathogenic and environmental strains (Powell and Loutit, 2004).

In recent years, the use of PCR-related tools has gained wide acceptance in developing countries. The advent of PCR has led to important advances in the development of routine diagnostic tests, and it has been possible to develop probes aimed at the detection of pathogen genetic material in host tissue, as well as for assessing genetic variability within and between fish and shellfish populations. Both DNA- and RNA-based methods have been devised to detect pathogen genetic material. Depending on the pathogen, conventional PCR can be replaced by the more sensitive nested PCR method, in which primers within the region amplified in a first step are used for further amplification of DNA. RNA quantification can be carried out using RT-PCR of the viral nucleic acids present in sample tissues. As with the immunological methods described earlier, it should be noted that PCR does not demonstrate the presence of disease nor of a viable pathogen, but only that pathogen genetic material was present in the sample being examined. Despite this limitation, and other problems related to ease of contamination, false positives, the limited number of primers available, etc., when properly applied, PCR offers a relatively rapid and inexpensive way for the routine screening of large numbers of aquatic animals for commercial aquaculture and for testing of imported stocks during quarantine. For example, PCR is very important in the routine screening of massive numbers of penaeid shrimp larvae for serious viral pathogens such as WSSV, TSV etc. in Asian and Latin America countries.

DNA probes and epidemiology

DNA probes have particular value in field epidemiology, routine disease surveillance and monitoring, treatment and eradication programmes in aquaculture and efforts to prevent the spread of pathogens to new geographical areas. These biotechnologies also have important application in risk management for aquatic animal diseases, including inspection and certification of production, and facilities and consignments for freedom from specific pathogens, achieving recognition of a country as having disease-free status, implementing disease zoning programmes, implementing effective quarantine measures, etc. (Bernoth, 2008).

The Manual of Diagnostic Tests for Aquatic Animals, regularly published by the World Animal Health Organisation (OIE) validates the use of traditional diagnostic methods such as evaluation of clinical signs, necropsy, histopathology, parasitology, bacteriology, virology, mycology etc., as well as immunological tests such as ELISA for the presumptive and confirmatory identification of OIE-listed diseases. The introduction to the Manual notes that "molecular methods for fish diseases are recommended for either direct detection of the pathogen in clinically diseased fish or for the confirmatory identification of a disease agent isolated using the traditional method. With one or two exceptions, molecular techniques are currently not acceptable as screening methods to demonstrate the absence of a specific disease agent in a fish population for the purpose of health certification in connection with international trade of live fish and/or their products. There is a need for more validation of molecular methods for this purpose before they can be recommended in the Aquatic Manual". (OIE, 2009; see also Adams and Thompson, 2008). This highlights the importance of further validating these diagnostic tools for serious and emerging diseases across a range of different laboratories worldwide.

3.2.2 Vaccines

Adams *et al.* (2008) reviewed the vaccine technologies in aquaculture. Vaccination is the action in which a host organism is exposed to organic (biological) molecules that allow the host to mount a specific immune reaction through which it has a better capability to fight subsequent infections of a specific pathogen compared with genetically similar non-vaccinated hosts. It has also been shown to be cost-effective and has led to the reduction in use of antibiotics. In Norway, for example, antibiotic use has decreased from 47 tons to approximately one ton annually (Markestad and Grave, 1997 and Figure 4).

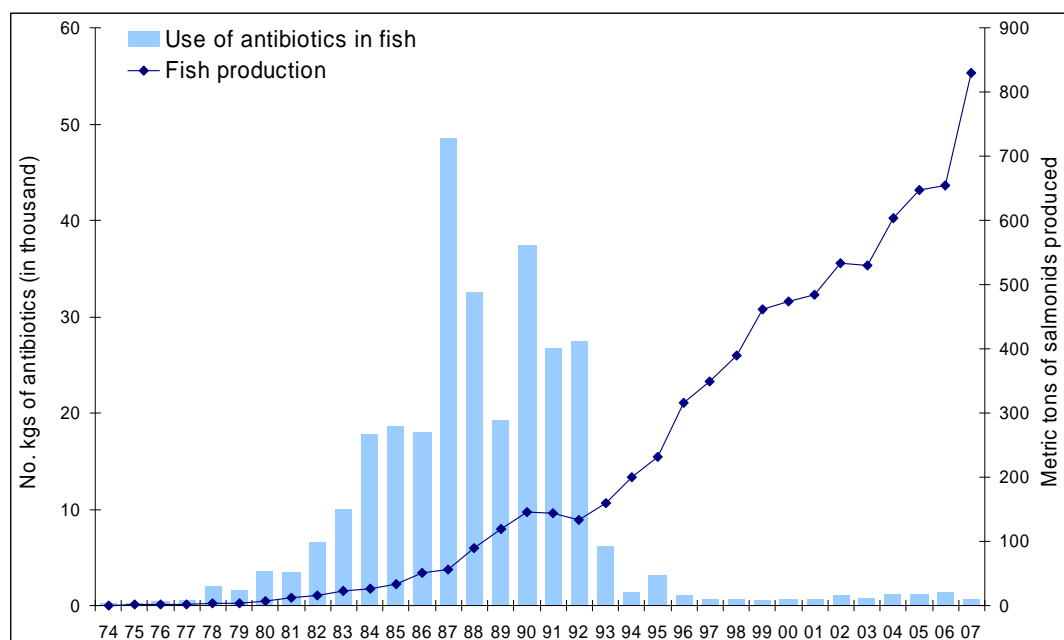
A wide range of commercial vaccines is available against bacterial and viral pathogens and many new vaccines are under development. Most target salmon and trout, and there are expanding opportunities for marine fish (Thompson and Adams, 2004). Traditionally, the organic molecules used for vaccination are directly derived from the pathogen in question. The most straightforward approach is to culture the pathogen after it has been inactivated and presented to the host. So far, vaccines containing more than ten bacterial pathogens and five viral pathogens have been produced based on such inactivated antigens (Sommerset *et al.*, 2005). Alternatively, the pathogen is not inactivated but chemically or genetically weakened so as to survive only for a limited period in the host, where it induces a specific immune response without causing disease and mortality. Such vaccines are generally described as "live" vaccines, and there is concern that the attenuated strain may back-mutate and revert to the virulent wild type (Benmansour and de Kinkelin, 1997). Due to environmental and control concerns in most countries, only two live bacterial (*Edwardsiella ictaluri* and *Flexibacter columnarum* for Channel catfish in the USA) and one live viral vaccine (koi herpesvirus for carp, in Israel) are commercially available at present.

One of the most important factors leading to reduced antibiotic use by the aquaculture sector is the availability of good prophylactic measures for diseases causing severe mortalities in cultured fish and shellfish. The use of vaccines provides good immunoprophylaxis for some of most important infectious diseases of finfish. In developed countries, their use has proved very effective at decreasing the unsustainable use of antibiotics. For example, in Norway antibiotic use in salmon farming has become almost negligible, at less than 1g per tonne of production, due mainly to the availability of vaccines for furunculosis and cold water vibriosis (Figure 4) (Smith, 2008). At almost similar production levels, Chilean salmon farming shows much more antibiotic

use due to the emergence of *Piscirickettsia salmonis*, a pathogen causing severe disease losses prior to harvest. Thus there have been recent attempts to develop immunoprophylactic measures.

As molecular-based vaccine production procedures rely heavily on biotechnological tools, vaccines are produced mainly in developed countries. A DNA vaccine is a circular DNA plasmid that contains a gene for a protective antigenic protein from a pathogen of interest (Kurath, 2008). Considerable industrial research has been conducted towards developing DNA vaccines for species such as salmonids for pathogens (generally, viruses) for which traditional methods have not been successful. As many strains and varieties of a single pathogen are generally present in the tropics, unlikely in temperate pathogens, monovalent vaccines are not practical under tropical conditions. Such difficulties, together with the lack of adequate biotechnological knowledge and financial resources, have led to slighter advances in vaccine development in the tropics, and for tropical species. Commercial vaccines using inactivated bacterial pathogens are available for some species: channel catfish, European seabass and seabream, Japanese amberjack and yellowtail, tilapia, Atlantic cod, salmon and trout (Ingunn *et al.*, 2005). Fewer commercially available viral vaccines have been produced, and no commercially available parasite vaccines exist.

Figure 4. Use of antibiotics vs. production of fish in Norway from 1974 to 2008 (Prof. Tore Hastein personal communication)



3.3 Environmental management and bioremediation

Aquaculture, like any other live production system, produces effluents rich in nutrients. Some aquatic production systems also produce effluents with harmful substances such as residues and metabolites of antibacterials and therapeutics. Developing systems that produce effluents with acceptable standards and improving the quality of the aquatic environment where effluent discharges are unacceptably high is a challenge. Biotechnological interventions such as bioremediation, the use of probiotics and vaccination offer significant promise for addressing these important issues.

Bioremediation is a promising biotechnological approach for the degradation of hazardous waste to environmentally safe levels using aquatic micro-organisms, or other filtering macro-organisms. Although this procedure has been used in various situations, such as sewage treatment (e.g. FAO, 2008), application to shrimp and other aquaculture wastes is fairly novel. There are a lot of commercial products on the market, mainly bacterial preparations, but the mode of action and

efficacy of many of these have yet to be scientifically measured. In addition to microbes, bivalves, seaweeds, holothurians (sea cucumbers), etc., have been tested to assess their ability to reduce organic loading, or reduce the excess nutrients produced during culture production. Various bioremediation preparations have also been developed with a view to removing nitrogenous and other organic waste in water and bottom sludge and thus reduce chemically-induced physiological stress, e.g., in pond-reared shrimp. More products will undoubtedly emerge with continued research in this field, but controlled field trials are urgently needed to determine the cost-benefit and effectiveness of these products under culture conditions.

Probiotics are generally administered as live microbial feed supplements which affect the host animal by improving the intestinal microbial balance to optimize the presence of non-toxic species. A stable gut microflora helps the host resist pathogenic invasions, particularly via the gastro-intestinal tract. Antibiotics reduce specific or broad-spectrum gut microflora and probiotics may have post-antibiotic treatment potential for restoring the microbial balance. Probiotics are widely used in animal husbandry but their use in aquaculture is still relatively new. However, there are increasing reports of potential probiotics for shrimp aquaculture, which has been plagued by opportunistic bacteria, such as the luminescent *Vibrio harveyi*, and, in some cases, probiotics have been reported to significantly reduce antibiotic use in shrimp hatcheries. Suppression of proliferation of certain pathogenic bacteria (e.g., *Vibrio* spp.) in shrimp hatcheries has been achieved by introducing (inoculating) non-pathogenic strains or species of bacteria that compete for microbial metabolite resources. This procedure shows promise to be effective and economical. However, further refinement of administration and concentration loads required for effective pathogen suppression is required. Effective and economically viable probiotics also require greater research into optimal strains of probiotic micro-organisms and stringent evaluation under field conditions of their economic feasibility.

As discussed earlier, the control of disease using vaccines is a reputed technology. There are interesting examples of reducing antibacterial use in aquaculture through the use of vaccination, particularly in temperate species such as salmon and trout. Reduction of the use of antibacterials not only diminishes the risk of rejection of aquatic products at international trading borders due to the presence/detection of residues above acceptable levels, it will also help in reducing the contamination of natural water bodies with harmful residues and the development of antimicrobial-resistant bacteria.

The proliferation of red tides with the blooming of harmful algae has been increasingly reported in many parts of Latin America, where the toxins represent a threat to food safety as well as a cause of fish and shellfish losses from the associated mortalities. Red tides can produce significant economic losses to fisheries and aquaculture due to bans on the marketing of fish and shellfish from the affected geographical area and to the toxic effects on fish. In Central America and the Caribbean, la “ciguatera” is the most important cause of toxic poisoning resulting from consumption of tropical fish. In Latin America, blooms of *Alexandrium* spp. are one of the major causes of large economic losses due to the banning of commercial sales of mussels. In Chile, preventive closures cause about US\$100 million in annual losses to the artisanal bivalve fishery. Furthermore, these closures have a direct negative impact on local employment in the shellfish production sector, which is labour intensive, thus having a detrimental effect on livelihoods. While it is not known if climate change is increasing the number of episodes of algal blooms, it is recognized that red-tide episodes have recently become more common. (Jessup *et al.*, 2009). Warm episodic currents also play a key role in causing large economic losses through mass mortalities of fish (Kedong *et al.*, 1999).

To date, the detection of toxins due to algal blooms is carried out using mouse bioassays and high performance liquid chromatography, but new methodologies are being developed for detection of *Alexandrium catenella* (Uribe and Espejo, 2003). Expressed sequence tag (EST) libraries are now publicly available (Uribe *et al.*, 2008), so that it may be possible to develop molecular diagnostic techniques. To improve the prevention of impacts on aquaculture, PCR techniques and EST libraries can be used also to assist the early detection of toxin-producing algae in vast marine areas.

3.4 Biodiversity conservation and fisheries management

Restocking procedures are a common practice in many developing countries, but the potential of restocking and stock enhancement stems primarily from the development of the technologies used to produce hatchery-reared juveniles (Bell *et al.*, 2006). The production of large numbers of juveniles and their subsequent release into the wild can affect a fishery resource in at least two ways (Bell *et al.*, 2006): (i) when stocking is done to restore a spawning biomass, there is some scope for interbreeding between the natural population and the introgressed population, and (ii) there might be enough individuals used to restore the carrying capacity of the fishery.

From a genetic point of view, the main consequence of restocking may be the hybridization of non-native individuals with natural stocks, which can have important impacts on natural biodiversity. Fish are very prolific, and under many hatchery production systems a relatively small number of parents can provide sufficient numbers of juveniles for release, in which case, the genetic variability of the fishery may be reduced. This situation can easily lead to genetic bottlenecks, the forthcoming generations of population being subjected to relatively high rates of inbreeding, thus inadvertently reducing the genetic variability of the population (Povh *et al.*, 2008). This can have large effects on the sensitivity of individuals to environmental variations, and could possibly cause the extinction of a population or species in a particular environment (Guttman and Berg, 1998). In addition, inbreeding can affect growth and reproduction. The mating of wild fish with those released by restocking programmes can promote the loss of genes important for local adaptation (Vasemägi *et al.*, 2005; Sønstebo *et al.*, 2007) in a genetic mechanism called out-breeding depression. While this concern has been effectively studied in terrestrial animals and in salmon populations in developed countries, this is not the case in other fisheries from developing countries. Therefore, careful restocking procedures need to be developed in order to reduce the potential for the introgressed population to reduce the genetic variability and therefore the sustainability of the resource. Assessing the genetic diversity of managed stocks or highly selected populations is an important issue when pedigree information is lacking or in situations where some kind of quality assurance is needed.

The use of molecular markers and the principles of population genetics have proved very effective in assessing the actual levels of genetic variability within single populations and in measuring the extent of differentiation between populations. For example, the Centro Nacional de Pesquisa de Peixes Tropicais in Brazil has studied the use of RAPD markers for the Amazonian fish “matrinxa” (*Brycon cephalus*) and has shown a relatively large reduction in genetic variability in fish used for restocking purposes compared with the native Amazonian river population (Povh, 2008).

In developing countries, the markers have been used mainly for assessing genetic variation in tilapia and carp populations in Thailand, the Philippines and India. Markers have been used for characterizing stocks and comparing levels of genetic variability in *Oreochromis* species. Agustin (1999) used markers to assess genetic differences between indigenous samples from Africa and populations from Asia, concluding that the low performance of *O. mossambicus* stocks can be explained by the effect of large bottlenecks in the populations used for aquaculture in Asia. Molecular markers have also been used to assess population differentiation of Nile tilapia (*O. niloticus*) for both domesticated and feral populations (Agnese *et al.*, 1997). In both cases, moderate to great genetic differentiation was found between strains, and the use of markers successfully correlates with the actual biogeographical data.

The escape of farmed fish from aquaculture may influence the genetic variability of native populations. The possible genetic impacts resulting from introductions and invasive alien species include: interbreeding between alien and native genotypes causing, in some cases, reduced reproductive efficiency generating nonviable offspring; decreased fitness from loss of co-adapted gene complexes; and indirect genetic impacts resulting from other ecological interactions (FAO, 2005a).

Climate change and related climatic events such as the El Niño-Southern Oscillation (ENSO) can have serious impacts on the distribution of fishery resources between countries. Based on census

numbers, mackerel fisheries were apparently depleted in Chilean coastal waters during the occurrence of ENSO episodes. However, markers have shown little differentiation with other populations in the Pacific Ocean (such those observed in New Zealand), and so it is likely that the drop in numbers is related to migration of the mackerel populations to colder waters in the Pacific rather than to fishery depletion (IFOP, 1996).

3.5 Concluding remarks

Aquaculture, compared with livestock and crop production, is a novel production system in many developing and developed countries. As shown in this section, biotechnologies are being applied in fisheries management, but their use is very limited compared to aquaculture. The use of successful and effective biotechnologies in aquaculture is very much confined to genetic manipulations and improvements and to health management.

The success or failure in using biotechnologies in developing countries depends to a large extent on: i) the markets for each of the products within the production sectors, and ii) the investment and acquisition capacity for the fisheries and aquaculture sectors. In the case of aquaculture, the latter is very important considering that the largest proportion of world production comes from developing countries and from small farmers (specifically in Asia). Most biotechnological interventions have been developed for improved production and the better management of aquaculture. Most have been targeted towards high-value commercial aquaculture species generally produced for international markets. Although many small-scale farmers are producing for export markets, the significant uptake of many biotechnological interventions and innovations has generally been restricted to commercial or industrial aquaculture operations. This is certainly due to the cost of the technology as well as the organized nature of industrial aquaculture.

Recently, however, as a result of better organization in the small-scale farming sector, certain biotechnologies have been effectively taken up by the small farmers in many parts of the developing world. They include DNA probes for detecting pathogens in some species (mainly PCR detection of major viral pathogens of shrimp), the use of SPF shrimp broodstock or postlarvae, the use of certain DNA vaccines, the all-male (genetically male) tilapia and, in some cases, markers for pedigree evaluation in salmon are being successfully introduced in small-scale production systems (see case studies) worldwide. In fact, almost everywhere in the world, shrimp farmers, whether small or large, currently use only PCR-tested postlarvae for stocking. For example, in India there are more than 90 laboratories providing PCR services for the shrimp sector – mainly the screening of seed and broodstock. In Vietnam there are over 40 laboratories. This pattern holds true in many countries of the region as the cost of using such biotechnologies has declined over the years and the benefit increased tremendously.

As mentioned above, the majority of aquaculture produce comes from the small-scale farming sector, in many instances comprising low-input extensive production systems. Although there is scope for biotechnologies, and although they are already being employed by small-scale farmers, classical environmental improvements and better management practices, such as conventional genetic selection of broodstock, conventional health management through the avoidance of pathogens, etc., can also contribute significantly towards improving small-scale aquaculture production and sustainability.

4. Case studies

Biotechnologies are used in aquaculture for reducing losses due to diseases and improving production through genetic manipulation. These technologies are regularly used in almost all countries, at different rates and levels, based on the intensity and commerciality of the production system. Here, two case studies are presented, outlining specific successful applications of biotechnological tools in aquaculture in developing countries.

4.1 PCR-based pathogen detection in shrimp aquaculture in India

At present, shrimp is the most valuable aquaculture commodity sector in the world. This sector has been continuously facing the challenge of new diseases, particularly viral pathogens. Some 20 years ago, there was hardly any accurate molecular-based pathogen detection system available in any part of the world. Now, however, as a result of advanced molecular research and biotechnology, there are many DNA-based detection technologies such as PCR methodologies available for all the major shrimp viruses. A number of PCR, nested-PCR and hybridization tests have been developed for virus detection. The tests use a range of different PCR primers and hybridization probes targeted to different and poorly defined sites in the virus genome. Several RT-PCR tests are also available. The application of PCR detection of viruses of broodstock and postlarvae in both *Penaeus monodon* and *Penaeus vannameii* is now practised in all countries producing commercial shrimp at all levels (Karunasagar and Karunasagar, 1999; Lo *et al.*, 1998; Peinado-Guevara and López-Meyer, 2006). Recently, lateral flow chromatographic immunodiagnostic strips similar to common drug-store pregnancy tests have begun to appear for some shrimp diseases. Using these, unskilled farm personnel can easily diagnose shrimp disease outbreaks at the farm. The strips are relatively cheap and quick (e.g. www.biotech.or.th/sbbu/ENG/index.asp). Other methods comparable to PCR and RT-PCR are now available or are being developed for single and dual or multiple viral detection, but they too currently require advanced equipment and personnel.

This rapid-detection technology has given a new dimension to the shrimp industry and losses due to viral diseases in shrimp have been reduced tremendously by the use of PCR-tested postlarvae for stocking. Recent successes in farmer group or cluster formation and management in shrimp aquaculture, particularly in India and Indonesia, are to a large extent based on good health management which includes the use of PCR tested postlarvae for stocking in ponds. This demonstrates a scenario in which a successful biotechnology has not only contributed towards realizing its scientific objective, but also towards improving the overall governance of the sector (Subasinghe *et al.*, 2008).

If a specific situation is considered, the use of PCR detection technology was the key, basic step towards developing an effective better management practice (BMP) for small-scale shrimp aquaculture in Andhra Pradesh. In India, aquaculture is mainly carried out by small and marginal scale farmers located in the remote villages of the country. They are largely unorganized, scattered and poorly educated. The farmers mostly opt for traditional methods for operating their farms and do not have access to technological innovations or scientific applications. A joint MPEDA-NACA (Marine Products Export Development Authority – Network of Aquaculture Centres in Asia-Pacific) project assisted by FAO to support shrimp farmers in disease control and coastal management was initiated in 2002, leading to the participatory development of BMPs that provided significant improvements in profits and reduced shrimp disease risks for farmers. One of the key interventions that the farmers adopted in applying BMPs in their quest to reduce losses due to disease was the use of PCR-screened postlarvae for stocking.

The project supported farmers in the implementation of BMPs through the formation of self-help groups around local “clusters”. An economic analysis of 15 farmer groups in Andhra Pradesh clearly demonstrated that farmers adopting BMPs, including the use of PCR-screened postlarvae for stocking, had higher profitability, lower production costs and were able to produce quality and traceable shrimp without using any banned chemicals.

The project has been highly successful in forming a self-help movement of farmers across India through a grass-roots approach. From a mere five farmers who first adopted the cluster-farm approach and BMPs in 2002, the programme had swelled to more than 1 000 farmers in 30 aquaculture societies in five coastal states by 2007. Beginning in 2007, the MPEDA-NACA project became the National Centre for Sustainable Aquaculture (NaCSA). NaCSA is an outreach organization of MPEDA established to service the small-scale aquaculture sector and provide technical support to farmer groups. It aims to empower and build the capacity of small-scale farmers to produce quality shrimps in a sustainable and more profitable manner.

Perhaps one of the keys to the above success is the ability to reduce losses due to disease in production systems, and to a large extent this has been possible through the use of PCR technology for screening and detecting major viral pathogens in broodstock and postlarvae.

4.2 Specific pathogen-free (SPF) stocks in shrimp aquaculture

Only a few species have so far been domesticated in the aquaculture sector. One group of species on which most research has been focused on the domestication and development of SPF strains is the penaeid shrimp. SPF shrimp are produced in SPF facilities using many biotechnological tools, particularly DNA-based pathogen detection and diagnostic techniques. The primary goal of SPF facilities is to produce strains of shrimp that are disease-free, domesticated and genetically improved for aquaculture. SPF lines are available for *P. vannamei*, *P. stylirostris* and *P. monodon*. The SPF status should signify that the shrimp have passed through a rigorous quarantine and disease-screening process that has found them to be free from specified pathogens of concern to culturists. This characteristic means that countries or regions which still do not have this species can be reasonably sure that the importation of SPF animals will not result in the introduction of the specified pathogens from which the animal is declared free. This does not, however, guarantee against the animal being infected with unknown pathogens or known pathogens that are not screened against.

Genuine SPF shrimp are produced in biosecure facilities that have been repeatedly examined and found free of specified pathogens using intensive surveillance protocols, and originate from broodstock developed with strict founder population development protocols. These founder populations are generated by extensive quarantine procedures that result in SPF F1 generations derived from wild parents. Only stocks raised and held under these conditions, can be considered truly SPF. There is not yet an internationally agreed protocol for the development of SPF shrimp, and certainly some variation in the quality of different SPF stocks exists. Once the animals are removed from the SPF production facilities, they should no longer be referred to as SPF, even though they may remain pathogen-free. Once outside the SPF facility, the shrimp may be designated as High Health (since they are now subject to a greater risk of infection), but only if they are placed into a well-established facility with a history of disease surveillance and biosecurity protocols. If the shrimp are put anywhere else, for example into a non-biosecure maturation unit, hatchery or farm, they can no longer be called SPF or High Health as they are now exposed to a high risk of infection (FAO, 2005b).

One potential drawback of SPF animals is that they are only SPF for the specific diseases for which they have been checked. Typically this will consist of the viral pathogens which are known to cause major losses to the shrimp culture industry, including WSSV, YHV, TSV, IHHNV, *Baculovirus penaei* Virus and Hepatopancreatic Parvovirus as well as microsporidians, haplosporidians, gregarines, nematodes and cestodes. Despite this screening, new, hidden or "cryptic" viruses may be present, but because they are as yet unrecognized, may escape detection. Thus, it is believed that SPF shrimp shipped from Hawaii resulted in the contamination of shrimp in Brazil and Colombia with TSV. This was because, at the time, TSV was not known to have a viral cause and therefore went unchecked in SPF protocols.

In any case, the use of SPF stocks is only one part of a complete plan for minimizing disease risks in shrimp culture. The development of SPF strains is really designed to ensure that postlarvae stocked into grow-out ponds are free of disease, which is one of, if not the most serious, sources

of contamination. Other areas of this strategy that must be implemented include: strategies to ensure broodstock, eggs, nauplius, larvae and juveniles derived from SPF stock remain SPF.

Creating an enabling public sector environment is essential to better governance at all levels of aquaculture development. There have been many regulatory rebounds in aquaculture sector, in particular in shrimp farming in some countries. Uncontrolled and unregulated development of the sector has outstripped the carrying capacity in some locations, causing significant production losses mainly due to disease and resulting in the complete abandonment of farms. Significant improvements have been made in mitigating such catastrophic problems, and the negative environmental and social impacts of shrimp farming throughout the world have been significantly reduced. The use of wild-caught postlarvae in shrimp culture, which has a significant impact on aquatic biodiversity, has almost stopped or is little practised. The recent development of SPF broodstocks of some species of shrimp has reduced reliance on wild-caught postlarvae to a minimum.

SPF shrimp if produced and maintained under good biosecurity have proved successful. The success of SPF stocks may be more pronounced in large-scale industrial shrimp culture facilities where maintaining stringent biosecurity is possible. The use of this successful biotechnological approach in the rather disorganized small-scale shrimp aquaculture production sector poses another challenge (Briggs, Subasinghe and Funge-Smith, 2004).

B. Looking forward: Preparing for the Future

5. Key issues where biotechnologies could be useful

5.1 Environmental sustainability

Aquaculture is the fastest growing food producing sector in the world. It is poised to expand, diversify and intensify over the coming decades to bridge the increasing global gap between the supply of and demand for aquatic food. Responsible production through sustainable practices is the key to achieving this massive task. In the effort to maximize the contribution from aquaculture, it is inevitable that many constraints and hurdles need to be overcome. The biggest hurdle is to maintain environmental sustainability,

Conventional methods of controlling diseases, such as chemotherapeutics, are ineffective for many new pathogens (notably viruses). Molecular techniques have therefore received increasing attention for pathogen-screening and identification. In addition, these biotechnologies are providing significant insights into pathogenesis (disease development) and show strong potential for disease control and prevention programmes (e.g. DNA vaccines), as well as for treatments of diseases. The increased sensitivity and specificity conferred by DNA- or RNA-based probes has provided significant inroads for the early detection of diseases and identification of subclinical carriers of infections. This has had a direct effect on enhancing preventative management and control of disease in cultured species. Concomitant with this has been a decrease in the need for reactive treatments using traditional methodologies such as antibiotics or culling and disinfection. This has been particularly successful for shrimp broodstock selection and has broken the infection cycle perpetuated for years by accidental broodstock transmission of viral pathogens to developing offspring.

Biotechnologies can provide much assistance to better aquatic animal health management in aquaculture in developing countries, in particular through the development of sensitive and accurate molecular diagnostic methods and tools as well as vaccines for tropical diseases. Bioremediation and probiotics also provide some further opportunities.

5.2 Climate change

In the future, one of the greatest constraints could be the impact of climate change on aquaculture. Climate change threatens fisheries and aquaculture through higher temperatures and changes in weather patterns, water quality and supply. Important differences in the magnitude and types of impacts on aquaculture are predicted for different regions. The ability to adapt will confer a major advantage and should be developed by countries and regions. There is a need for the aquaculture sector to join other economic sectors in preparing to address the potential impact of global warming. One of the practical responses to climate change for aquaculture could be to strengthen the adaptive capacity and resilience of the sector, particularly that of small farmers and aquatic resources users. Increased resilience is a desirable feature of any sector; it can mitigate the future impact of unforeseen events (e.g. economic change, disease epidemics, Tsunamis, etc.), including those related to climate. There is some knowledge and experience from aquaculture itself and from the broader area of agriculture and natural resource management which could be used. Aquaculture, and particularly mariculture, could in fact provide adaptation opportunities to produce good quality protein when freshwater may become scarce. On the other hand, freshwater aquaculture can produce protein with higher water saving than other animal production sectors. Certain biotechnologies, particularly those dealing with genetic improvement, health and environmental mitigation should be of significant value for the discovery of adaptive technologies and interventions to counter the ever-present menace of climate change.

6. Identifying options for developing countries

Aquaculture is the fastest growing food-producing sector in the world. To bridge the future gap between demand for and the supply of aquatic food simply to maintain the current level of consumption, production needs to be almost doubled in less than three decades. In the quest to meet this unprecedented demand in the coming years, the aquaculture sector will face serious constraints. Four major constraints are inevitable: a) disease prevention and health management, b) genetic improvement and domestication, c) environmental management and d) food safety. These constraints are not new. They have been constantly addressed during the development of aquaculture over the past two decades, including through the use of biotechnologies.

Over the years, aquaculture biotechnology and other technological innovations have had a positive impact on aquaculture diversification, investment potential, and international technology exchange. The development of biotechnology in aquaculture should therefore provide a means of producing healthy and fast-growing animals by environmentally friendly means. However, this development will largely depend on the desire and willingness of the producers to work hand-in-hand with scientists and on the international donor community's readiness to assist developing countries in the related research, capacity-building and infrastructure development. The improved exchange of information and discussion between scientists, researchers and producers from different regions about their problems and achievements will undoubtedly help this important sector to develop with a view to increasing sustainable global aquatic animal production.

Based on the overview and previous analyses contained in this document, a number of specific options can be identified for developing countries to help them make informed decisions regarding the adoption of biotechnologies in the future, such as when – and if – they should deploy one or more biotechnologies and, if they decide to do so, how they can ensure the successful application of the chosen biotechnologies to enhance food security in the future.

6.1 Analysis in the document shows that there are few biotechnological advancements and tools currently in use in small-scale aquaculture operations aiming at rural development, poverty alleviation and food security. However, there is a need to identify those, their application and socio-economic impact in developing countries.

Developing countries should therefore collect information on the aquatic animal biotechnologies that may be used and analyse their national-level adoption and the socio-economic impacts. Such information should be used to advise policy-makers on the cost/benefit implications of such application. Increased efforts should be made to develop aquatic biosecurity policies within national R&D programmes or national aquatic production programmes.

6.2 The use of biotechnologies in aquaculture worldwide has increased incrementally over the past two decades. Several aquaculture biotechnologies have been used for the benefit of improved aquatic food production in both developed and developing countries, and have significant potential for future improvement. Most aquaculture biotechnologies are still too technical and costly for small-scale farmers. Efforts should be made to develop low-cost simple technologies that are easy to introduce to less advanced small-scale aquaculture farmers. Developing countries should give priority to developing aquaculture biotechnologies which are appropriate and conducive for both industrial and small-scale farmers.

6.3 Major biotechnological achievements and advances in fisheries and aquaculture have been mainly restricted to aquaculture and to the fields of genetics, health and the environment. Genetic improvements using gene manipulation (diploidy, triploidy) and hormonal therapy, etc. have shown promise for producing fish and shellfish with improved and desirable production qualities. Disease prevention and health management in aquaculture have benefited significantly from advances in biotechnologies. Many reliable and accurate rapid diagnostic techniques which can be used by small-scale farmers have been developed. There are several efficient vaccines now available for certain aquaculture species, which has significantly reduced the use of antibacterials

in their culture. However, more research is required to develop vaccines for tropical species, particularly the major species of global production. Some environmental remediation tools and technologies have been developed using several biotechnologies. They are being applied in some production systems, but their broad adoption across different production systems and practices is yet to be established.

The potential contribution of biotechnologies for genetic improvement to improve production of culture aquatic species should be recognized. National research and development plans should include appropriate research in these areas. In aquatic animal health research, the development of molecular diagnostics, vaccines and probiotics should be prioritized and national research institutions should also carry out research using appropriate biotechnologies that can help development of sustainable aquaculture in this area. National governments embarking on aquaculture development should also recognize that there is ample evidence for positive aquatic environmental impacts using various biotechnological interventions, thus use of biotechnology for improving the aquatic environment should be considered.

6.4 Until recently, perhaps because the application of fisheries and aquaculture biotechnologies has been mainly restricted to the commercial and industrial aquaculture of temperate species, there has been little evidence in many developing countries of national-level efforts to prioritize the development and application of biotechnologies in aquaculture. Even when efforts to develop such biotechnologies in the public sector were made in developing countries, they were not always directed towards or made available to improve small-farmer livelihoods. There is a need to create national policy environments in developing countries, including suitable investment and funding opportunities, to allow the development and application of appropriate biotechnologies in support of aquaculture development. National governments should pay special attention to the small-scale aquaculture sector. Preferential treatment of the sector towards capacity building in appropriate biotechnologies should also be considered.

6.5 Required funding in developing countries for aquatic biotechnological research and applications should be found through national budgets or through extra budgetary resources. An integral part of funding should be directed towards investment in capacity-building in the relevant fields of the aquaculture sector. A suitable investment environment and funding opportunities should be created to allow the development and application of appropriate biotechnologies in support of aquaculture development. The appropriate involvement of the relevant stakeholders in decision-making processes should be assured.

6.6 The establishment of efficient institutional structures and enforceable legal frameworks are important for the responsible use of biotechnologies in aquaculture at the national level. Such institutional arrangements should also strengthen research and extension needs and enhance relevant human and infrastructural capacities. National legal frameworks in aquaculture biotechnologies should be developed within an integrated national biotechnology framework, which also complies with the legal or voluntary requirements of international treaties and agreements that the country has ratified.

6.7 National biotechnology programmes in developing countries should include a special committee to oversee the aquatic biotechnology programme and research. Such committees should be formed in all countries and regional cooperation should be sought.

6.8 Information gathering and dissemination on aquatic biotechnologies should be encouraged within and between countries in a given region, and developing countries should consider setting up dedicated websites for this purpose.

6.9 Aquaculture products are facing increasing competition in accessing international markets. One of the key criteria is food safety and compliance with international food safety standards. Many such standards can be met through better farming that uses both simple and advanced biotechnological interventions. The aquaculture industry should therefore consider the importance of such biotechnological interventions in improving and maintaining food safety of cultured aquatic products. National governments in developing countries should consider research and development interventions on food safety within the broader framework of biotechnology.

7. Identifying priorities for action for the international community

The international community, including FAO and other UN organizations, NGOs, donors and development agencies, can play a key role in supporting developing countries by providing a framework for international cooperation and funding support for the generation, adaptation and adoption of appropriate biotechnologies. Here, a set of Priorities for Action is identified that can assist the international community in playing this role.

7.1 Relevant international institutions, donors and development partners should recognize that biotechnological interventions can contribute to sustainable aquaculture development worldwide.

7.2 Relevant international agencies should assist developing countries to collect, collate and analyse information about the biotechnologies in use in fisheries and aquaculture, and their contributions to national food security, poverty alleviation and social development.

7.3 Relevant international agencies should make efforts to maintain databases and information systems to assist countries access information for national biotechnology development programmes relating to fisheries and aquaculture.

7.4 Donors and international funding agencies supporting sustainable aquaculture development for food security and poverty alleviation should dedicate an appropriate share of their assistance projects to promoting and strengthening aquatic biotechnology R&D in developing countries. International research efforts should focus on developing interventions that are accessible to small-scale farmers.

7.5 When supporting the application of biotechnologies in fisheries and aquaculture, the international community should consider that technical assistance in biotechnology R&D should not be done at the expense of funding for other key research fields, and that the technical assistance should support effective and intimate links to strong breeding and extension programmes.

7.6 The international community assisting developing countries towards aquaculture sustainability should consider biotechnological advancement as an important area to be supported, and should assist developing countries in strengthening capacities for biotechnology policy development and long-term planning.

7.7 The international community should assist developing countries to develop the capacities of the national agricultural research systems, which include aquaculture, to involve relevant stakeholders in decision-making processes.

7.8 The international community should assist developing countries in the development of adequate institutional capacities in the development and enforcement of regulations related to use of biotechnologies in fisheries and aquaculture.

C. References

- Adams, A. & Thompson, K.D.** 2006. Review: Biotechnology offers revolution to fish health management. *Trends in Biotechnology* 24: 201–205.
- Adams, A. & Thompson, K.D.** 2008. Recent applications of biotechnology to novel diagnostics for aquatic animals. *Rev. sci. tech. off. epiz.* 27:197–209. (Also available at www.oie.int/boutique/extrait/16adams197210.pdf).
- Adams, A., Aoki, T., Berthe, C.J., Grisez, L., & Karunasagar, I.** 2008. Recent technological advancements on aquatic animal health and their contributions toward reducing disease risks - a review. In M.G. Bondad-Reantaso, C.V. Mohan, M. Crumlish & R.P. Subasinghe, eds. *Diseases in Asian Aquaculture VI*. Manila, Philippines, Fish Health Section, Asian Fisheries Society.
- Agnèse, J.F., Adépo-Gourène, B., Abban, E.K. & Fermon, Y.** 1997. Genetic differentiation among natural populations of the Nile tilapia *Oreochromis niloticus* (Teleostei, Cichlidae). *Heredity* 79:88-96.
- Agustin, L.Q.** 1999. *Effects of population bottlenecks on levels of genetic diversity and patterns of differentiation in feral populations of Oreochromis mossambicus*. Ph.D. thesis, Queensland University of Technology, Brisbane, Australia.
- Aoki, T., Hirono, I., de Castro, T. & Kitao, T.** 1989. Rapid identification of *Vibrio anguillarum* by colony hybridization. *J. Appl. Ichthyol.* 5:67–73.
- Asknes, A., Gjerde, B. & Roald, S.** 1986. Biological chemical and organoleptic changes during maturation of farmed Atlantic salmon. *Aquaculture* 53:7–20.
- Basant K. Tiwary, R. Kirubakaran, A.K. Ray.** 2004. The biology of triploid fish. *Reviews in Fish Biology and Fisheries*, 14: 391-402.
- Basavaraju Y., Mair, G.C., Mohan Kumar, H.M., Pradeep Kumar, S., Keshavappa, G.Y. & Penman D.J.** 2002. An evaluation of triploidy as a potential solution to the problem of precocious sexual maturation in common carp, *Cyprinus carpio*, in Karnataka, India. *Aquaculture*, 204: 407–418.
- Bell, J.D., Bartley, D.M., Lorenzen, K & Loneragand, N.R.** 2006. Restocking and Stock Enhancement of Coastal Fisheries – Potential, Problems and Progress. *Fisheries Research* 80: 1–8.
- Benfey, T.J.** 1999. The physiology and behavior of triploid fish. *Res. Fish. Sci.*, 7:39–67.
- Benfey, T.J., Solar, I., De Jong, G., Donaldson, E.M.** 1986. Flow-cytometric confirmation of aneuploidy in sperm from triploid rainbow trout. *Transactions of the American Fisheries Society* 115:838–840.
- Benmansour, A. & de Kinkelin, P.** 1997. Live Fish Vaccines: History and Perspectives. In R. Gudding, A. Lillehaug, P.J. Midtlyng, & F. Brown, eds *Fish Vaccinol. Dev. Biol. Stand.* pp. 279–289. Basel, Karger.
- Bernoeth, E.** 2008. The role of OIE aquatic standards and OIE Reference Laboratories in aquatic animal disease prevention and control. *Rev. Sci. Tech. off. Int. Epiz.* 27: 39–54.
- Berthe, F., Bureson, E. & Hine, M.** 1999. Use of molecular tools for mollusc disease diagnosis. *Bull. Euro. Ass. Fish Patrol.* 19: 277–278.
- Bijma, P., van Sarandon, JAM. & Bovenhuis, J.H.** 1997. Breeding value and variance component estimation from data containing inbred individuals: application to gynogenetic families in common carp. (*Cyprinus carpio* L.). *Genetics* 145: 1243–1249.

Bondad-Reantaso, M.G. & Subasinghe, R. 2005. *Aquaculture Health International*, Issue 1, 4–5.

Bondad-Reantaso, M.G., Subasinghe, R.P., Arthur, J.R., Ogawa, K., Chinabut, S., Adlard, R., Tan, Z. & Shariff, M. 2005. Disease and health management in Asian aquaculture. *Vet. Parasitol.* 132, 249–272.

Briggs, M., Funge-Smith, S., Subasinghe, R. & Phillips, M. 2004. *Introductions and movement of Penaeus vannamei and Penaeus stylirostris in Asia and the Pacific*. Food and Agricultural Organization of the United Nations, Regional Office for Asia and the Pacific. RAP Publication 2004/10.

Brown, C.R., Woolliams, J.A. & McAndrew, B.J. 2005. Factors influencing effective population size in commercial populations of gilthead seabream, *Sparus aurata*. *Aquaculture* 247: 219–225.

Carnegie, R.B., Meyer, G.R., Blackbourn J., Cochenec-Laureau, N., Berthe, F.C.J. & Bower, S.M. 2003. Detection of the oyster parasite *Mikrocytos mackini* by PCR and fluorescent *in situ* hybridization, and a preliminary phylogenetic analysis using SSU rDNA. *Dis. Aquat. Org.* 54(3): 219–227.

Choubert, G., & Blanc, J. 1989. Dynamics of dietary canthaxanthin utilization in sexually maturing female rainbow trout (*Salmo gairdneri* Rich.) compared to triploids. *Aquaculture*, 83 359–366.

Cnaani, A., Hallerman, E.M., Ron, M., Weller, J.I., Indelman, M., Kashi, Y., Gall, G.A.E. & Hulata, G. 2003. Detection of a chromosomal region with two quantitative trait loci, affecting cold tolerance and fish size, in an F2 tilapia hybrid. *Aquaculture* 223: 117–128.

Cnaani, A., Zilberman, N., Tinman, S., Hulata, G. & Ron, M. 2004 Genome-scan analysis for quantitative trait loci in an F2 tilapia hybrid. *Mol. Genet. Genom.* 272: 62–172.

Cochennec, N., Le Roux, F., Berthe, F. & Gerard, A. 2000. Detection of *Bonamia ostreae* based on small subunit ribosomal probe. *J. Invertebr. Pathol.* 76: 26–32

Coimbra, M.R., Kobayashi, M.K., Koretsugu, S., Hasegawa, O., Ohara, E., Ozaki, A., Sakamoto, T., Naruse, K. & Okamoto, N. 2003. A genetic linkage map of the Japanese flounder, *Paralichthys olivaceus*. *Aquaculture* 220: 203–218.

Cowley, J.A., Dimmock, C.M., Wongteerasupaya, C., Boonsaeng, V., Panyim, S. & Walker, P.J. 1999. Yellow head virus from Thailand and gill-associated virus from Australia are closely related but distinct prawn viruses. *Dis. Aquat. Org.* 36: 153–157.

Cunningham, C.O. 2004. Use of molecular diagnostic tests in disease control: Making the leap from laboratory to field application. In Leung, K.-Y., ed. *Current trends in the study of bacterial and viral fish and shrimp diseases. Molecular Aspects of Fish and Marine Biology* 3: 292–312. World Scientific Publishing Co.

De Silva, S.S., Subasinghe, R.P., Bartley, D.M. & Lowther, A. 2004. *Tilapias as alien aquatics in Asia and the Pacific: A review*. FAO Fisheries Technical Paper. No. 453. Rome, FAO.

Doyle, W. & Herbinger C.M. 1994. The use of DNA fingerprinting for high-intensity, within-family selection in fish breeding. *Proceedings of the Fifth World Congress on Genetics Applied to Livestock Production* Vol. 19, pp. 364–37. Ontario, Canada, Department of Animal and Poultry Science, University of Guelph.

Eknath A.E. & Acosta B.O. 1998. *Genetic Improvement of Farmed Tilapias (GIFT) Project: final report, March 1988 to December 1997*. International Center for Living Aquatic Resources Management, Makati City, Philippines.

-
- Falconer, D.S. & Mackay, T.F.C.** 1996. *Introduction to quantitative genetics*, 4th edition. Harrow, Essex, U.K, Longman.
- FAO/NACA.** 2000. *The Asia Regional Technical Guidelines on Health Management for the Responsible Movement of Live Aquatic Animals and The Beijing Consensus and Implementation Strategy*. FAO Fisheries Technical Paper. No. 402. Rome, FAO.
- FAO.** 1995. *Selective breeding programmes or medium-sized fish farms* by D. Tave. FAO Fisheries Technical Paper Series 352. (Also available at www.fao.org/docrep/field/009/v8720e/V8720E00.htm).
- FAO.** 2005a. *Fisheries Issues. Impact of aquaculture on biodiversity*. Text by Devin Bartley, Heiner Naeve, Rohana Subasinghe. In: *FAO Fisheries and Aquaculture Department* [online]. Rome. Updated 27 May 2005. www.fao.org/fishery/topic/14853/en
- FAO.** 2005b. *Introductions and movement of two penaeid shrimp species in Asia and the Pacific*, by M. Briggs, S. Funge-Smith, R.P. Subasinghe and M. Phillips. FAO Fisheries technical paper 476. (Also available at www.fao.org/docrep/009/a0086e/A0086E00.htm)
- FAO.** 2007. *Marker-assisted selection: Current status and future perspectives in crops, livestock, forestry and fish*, by E. Guimaraes, J. Ruane, A. Sonnino, B.D. Scherf and J. Dargie (eds.). (Also available at www.fao.org/docrep/010/a1120e/a1120e00.htm)
- FAO.** 2008. *Coping with water scarcity: What role for biotechnologies?*, by J. Ruane, A. Sonnino, P. Steduto, and C. Deane. FAO Land and Water Discussion Paper No. 7. (Also available at www.fao.org/docrep/011/i0487e/i0487e00.htm)
- FAO.** 2009. *The state of world fisheries and Aquaculture 2008*. Fisheries and Aquaculture Department of the Food and Agriculture Organization (FAO) of the United Nations. Rome.
- FAO FishStat.** www.fao.org/fishery/statistics/programme/3,1,1/en
- Fitzsimmons, K.** 2001. Introduction to Tilapia sex-determination and sex reversal. Unpublished report (<http://ag.arizona.edu/azaqua/ista/reports/sexreverse.doc>).
- Galbreath, P. F. & Thorgaard, G. H.,** 1994. Viability and freshwater performance of Atlantic salmon (*Salmo salar*, brown trout (*Salmo trutta*) triploid hybrids. *Can. J. Fish. aquat. Sci.* 51: 16–24.
- Gjerde, B., Villanueva, B. & Bentsen, H.B.** 2002. Opportunities and challenges in designing sustainable fish breeding programs. *Proc. 7th World. Congr. Genetics Appl. Livestock Prod.* CD-ROM 06–01. Montpellier, France.
- Grant, W.S.** 2007. Status and trends in genetic resources of capture fisheries. In D.M.. Bartley, B.J. Harvey & R.S.V. Pullin, eds. *Workshop on Status and Trends in Aquatic Genetic Resources. A basis for international policy*. Rome, FAO.
- Guttman, S.I. & Berg, D.** 1998. Changes in the genetic diversity of aquatic organisms in the great lakes: causes and consequences. *Setae News* pp. 23–24.
- Hansen, M.M.** 2008. *The use of molecular markers for preserving genetic resources in wild fish populations*. Presentation at 'Biotechnology as a toolbox to study and monitor agricultural genetic resources'. FAO side event for the 13th meeting of the Subsidiary Body on Scientific, Technical and Technological Advice (SBSTTA) of the Convention on Biological Diversity, 22 February 2008. Rome, FAO headquarters. www.fao.org/biotech/docs/hansen.htm
- Hauser, L., Adcock, G.J., Smith, R.J., Ramirez, J.H. & Carvalho, G.R..** 2002. Loss of microsatellite diversity and low effective population size in an overexploited population of New Zealand snapper (*Pagrus auratus*). *PNAS* 99:11742–11747.

- Herbinger, C.M., O'Reilly, P.T., Doyle, R.W., Wright, J.M. & O'Flynn, F.** 1999. Early growth performance of Atlantic salmon full-sib families reared in single family tanks versus in mixed family tanks. *Aquaculture* 173:105–116.
- Houssay, B.A.** 1930. Acción sexual de la hipófisis en los peces y reptiles. *Revista de la Sociedad Argentina de Biología* 106:686–688.
- Houston, R.D., Haley, C.S., Hamilton, A., Guy, D.R., Tinch, A.E., Taggart, J.B., McAndrew, B.J & Bishop S.C.** 2008. Major Quantitative Trait Loci Affect Resistance to Infectious Pancreatic Necrosis in Atlantic Salmon (*Salmo salar*). *Genetics* Vol. 178, 1109–1115.
- Hulata, G., Wohlfarth, G. & Moav, R.** 1985. Genetic differences between the Chinese and European races of the common carp, *Cyprinus carpio* L. IV. Effects of sexual maturation on growth patterns. *Journal of the Fishery Biology* 26:95–103.
- Ingunn, S., Bjørn, K., Eirik, B. & Petter, F.** 2005. *Expert Review of Vaccines*, Volume 4, Number 1, February 2005, pp. 89–101 (13).
- IFOP.** 1996. *PROYECTO FIP N° 96–15: Migración de jurel desde y hacia la ZEE de Chile central*. Instituto de Fomento Pesquero.
- Jessup, D.A., Miller, M.A., Ryan, J.P., Nevins, H.M. & Kerkering, H.A.** 2009. Mass Stranding of Marine Birds Caused by a Surfactant-Producing Red Tide. *PLoS ONE* 4(2): e4550. doi:10.1371/journal.pone.0004550
- Karunasagar, I. & Karunasagar, I.** 1999. Diagnosis, treatment and prevention of microbial diseases of fish and shellfish. *Current Science* 76, 387–399.
- Kedong, P., Harrison, J., Chen, J. & Huang, W.** 1999. Red tides during spring 1998 in Hong Kong: is El Niño responsible? *Marine ecology progress series* 187: 289–294,
- Khoo, S.K., Ozaki, A., Nakamura, F., Arakawa, T., Ishimoto, S., Nickolov, R., Sakamoto, T., Akutsu, T., Mochizuki, M., Denda, I. & Okamoto, N.** 2004. Identification of a novel chromosomal region associated with infectious hematopoietic necrosis (IHN) resistance in rainbow trout. *Fish Path.* 39: 95–102.
- Kocher, T.D., Lee, W.J., Sobolewska, H., Penman, D. & McAndrew, B.** 1998. A genetic linkage map of a cichlid fish, the tilapia (*Oreochromis niloticus*). *Genetics* 148: 1225–1232.
- Kurath, G.** 2008. Biotechnology and DNA vaccines for aquatic animals. *Rev. Sci. Technol.* 27:175–196.
- Lallias, D., Lapegue, S., Hecquet, C., Boudry, P. & Beaumont, A.R.** 2007. AFLP-based genetic linkage maps of the blue mussel (*Mytilus edulis*). *Animal Genetics* 38(4):340–349.
- Lee, B.Y., Hulata, G. & Kocher, T.D.** 2004. Two unlinked loci controlling the sex of blue tilapia (*Oreochromis aureus*). *Heredity* 92, 543–549.
- Lee, B.Y., Lee, W.J., Streelman, J.T., Carleton, K.L., Howe, A.E., Hulata, G., Slettan, A., Stern, J.E., Terai, Y. & Kocher, T.D.** 2005. A second-generation genetic linkage map of tilapia (*Oreochromis* spp). *Genetics* 170: 237–244.
- Le Roux, F., Audemard, C., Barnaud, A. & Berthe, F.C.J.** 1999. DNA probes as potential tools for the detection of *Marteilia refringens*. *Marine Biotechnology* 1 (6): 588–597.
- Lightner, D.V.** 1996. *A Handbook of Shrimp Pathology and Diagnostic Procedures for Diseases of Cultured Penaeid Shrimp*. Baton Rouge, LA, USA, World Aquaculture Society.
- Lightner, D.V. & Redman, R.M.** 1998. Shrimp disease and current diagnostic methods. *Aquaculture* 164: 201–220.
- Liu, Z.J. & Cordes, J.F.** 2004. DNA marker technologies and their applications in aquaculture genetics. *Aquaculture* 238: 1–37.

- Lo, C.F., Chang, Y.S. & Chen, C.T.** 1998. PCR monitoring of cultured shrimp for white spot syndrome virus (WSSV) infection in growth ponds. In T.W. Flegel, ed. *Advances in Shrimp Biotechnology*, pp. 281–286. Bangkok, National Center for Genetic Engineering and Biotechnology.
- Mair, G.C.** 1999. *Genetic improvement of tilapias: application in aquaculture and future prospects*, pp. 86–107. The Fifth Roche Aquaculture Conference, Bangkok, August 26, 1999. Rovithai, Bangkok, Thailand.
- Markestad, A. & Grave, K.** 1997. Reduction of antibacterial drug use in Norwegian fish farming due to vaccination. In R. Gudding, A. Lillehaug, P.J. Midtlyng & F. Brown, eds. *Fish Vaccinol. Dev. Biol. Stand.* Basel, Karger.
- Martinez, V.** 2007. Marker-Assisted Selection in fish and shellfish breeding schemes. In E. Guimaraes, J. Ruane, A. Sonnino, B.D. Scherf & J. D. Dargie, eds. *Marker-assisted selection: Current status and future perspectives in crops, livestock, forestry and fish*, Chapter 17. Rome, FAO.
- Martinez, V., Kause, A., Mäntysaari, E. & Mäki-Tanila, A.** 2006. The use of alternative breeding schemes to enhance genetic improvement in rainbow trout: II. Two-stage selection. *Aquaculture* 254: 195–202.
- Martinez, V., Neira, R. & Gall, G.A.E.** 1999. Estimation of genetic parameters from pedigreed populations: lessons from analysis of alevin weight in coho salmon (*O. kisutch*). *Aquaculture* 330: 22–30.
- McGeachy, S.A., Benfey, T. J. & Friars, G. W.** 1995. Freshwater performance of triploid Atlantic salmon (*Salmo salar*) in New Brunswick aquaculture. *Aquaculture* 137: 333–341.
- Moen, T., Agresti, J.J., Cnaani, A., Moses, H., Famula, T.R., Hulata, G., Gall, G.A.E. & May, B.** 2003. A genome scan of a four-way tilapia cross supports the existence of a quantitative trait locus for cold tolerance on linkage group 23. *Aquaculture Research* 35, 893–904.
- Morris, D.C., Morris, D.J. & Adams, A.** 2002. Development of improved PCR to prevent false positives and false negatives in the detection of *Tetracapsula bryosalmonae*, the causative agent of Proliferative Kidney Disease. *J. Fish Dis.* 25(8): 483–490.
- Nichols, K.M., Young, W.P., Danzmann, R.G., Robison, B.D., Rexroad, C., Noakes, M., Phillips, R.B., Bentzen, P., Spies I., Knudsen, K., Allendorf, F.W., Cunningham, B.M., Brunelli, J., Zhang, H., Ristow, S., Drew, R., Brown, K.H., Wheeler, P.A. & Thorgaard, G.H.** 2003. A consolidated genetic linkage map for rainbow trout (*Oncorhynchus mykiss*). *Anim. Genet.* 34: 102–115.
- Norris, A.T., Bradley, D.G. & Cunningham, E.P.** 2000. Parentage and relatedness determination in farmed Atlantic salmon (*Salmo salar*) using microsatellite markers. *Aquaculture* 182: 73–83.
- OIE.** 2009. *Manual of diagnostic tests for aquatic animals*. Summary. (Also available at www.oie.int/eng/normes/fmanual/A_summry.htm)
- Overturf, K., LaPatra, S. and Powell, M.** 2001. Real-time PCR for the quantitative analysis of IHNV in salmonids. *J. Fish Dis.* 24: 325–333.
- Ozaki, A., Sakamoto, T., Khoo, S., Nakamura, K., Coimbra, M.R., Akutsu, T. & Okamoto, N.** 2001. Quantitative trait loci (QTLs) associated with resistance/susceptibility to infectious pancreatic necrosis virus (IPNV) in rainbow trout (*Oncorhynchus mykiss*). *Mol. Genet. Genom.* 265: 23–31.
- Paterson, S., Piertney, S.B. & Knox, D.** 2004. Characterization and PCR multiplexing of novel highly variable tetranucleotide Atlantic salmon (*Salmo salar* L.) microsatellites. *Mol. Ecol. Notes* 4: 160–162.

- Peinado–Guevara L.I. & López–Meyer, M.** 2006. Detailed monitoring of white spot syndrome virus (WSSV) in shrimp commercial ponds in Sinaloa, Mexico by nested PCR. *Aquaculture* 25: 33–45.
- Phelps, R.P.** 2001. Sex reversal: The directed control of gonadal development in tilapia. In D.E. Meyer, ed. *Proc. 6to. Simposio Centroamericano de Acuicultura*, pp. 35-60. Asociación Nacional de Acuicultores de Honduras and the Pond Dynamics/Aquaculture Collaborative Research Support Program. Tegucigalpa, Honduras..
- Ponzoni, R.W.N., Nguyen, H. & Khaw, H.L. Ling.** 2006. *Importance and implementation of simple and advanced selective breeding programs for aquaculture species in developing countries*. 8th World Congress on Genetics Applied to Livestock Production, August 13–18, 2006, Belo Horizonte, MG, Brazil.
- Povh, J.A., Lopera–Barrero, N.M., Ribeiro, R.P., Lupchinski Jr, E., Gomes, P.C. & Lopes, T.S.** 2008. Genetic monitoring of fish repopulation programs using molecular markers. *Cien. Inv. Agr.* 35(1): 5–15.
- Powell, J.L. & Loutit, M.W.** 2004. Development of a DNA probe using differential hybridization to detect the fish pathogen *Vibrio anguillarum*. *Microbial Ecol.* 28: 365–373.
- Primmer, C.** 2006. Genetic characterization of populations and its use in conservation decision-making in fish. In J. Ruane & A. Sonnino, eds. *The role of biotechnology in exploring and protecting agricultural genetic resources*. Rome, FAO. (Available at www.fao.org/docrep/009/a0399e/a0399e00.htm).
- Purdom, C.E.** 1983. Genetic engineering by the manipulation of chromosomes. *Aquaculture* 33: 287–300.
- Puttinaowarat, S., Thompson, K.D. and Adams, A.** 2000. Mycobacteriosis: detection and identification of aquatic Mycobacterium species. *Fish Vet. J.* 5:6–21.
- Reid, D.P., Szanto, A., Glebe, B., Danzmann, R.G. & Ferguson, M.M.** 2005. QTL for body weight and condition factor in Atlantic salmon (*Salmo salar*): comparative analysis with rainbow trout (*Oncorhynchus mykiss*) and Arctic charr (*Salvelinus alpinus*). *Hered.* 94: 166–172.
- Rodriguez, F.M., LaPatra, S., Williams, S., Famula, T. & May, B.** 2004. Genetic markers associated with resistance to infectious hematopoietic necrosis in rainbow and steelhead trout (*Oncorhynchus mykiss*) backcrosses. *Aquaculture* 241: 93–115.
- Routray, P., Verma, D. K., Sarkar S. & Sarangi N.** 2007. Recent advances in carp seed production and milt cryopreservation. *Fish Physiology and Biochemistry* 33: 413–427.
- Sakamoto, T., Danzmann, R.G., Gharbi, K., Howard, P., Ozaki, A., Khoo, S.K., Woram, R.A., Okamoto, N., Ferguson, M.M., Holm, L.-E., Guyomard, R. & Hoyheim, B.** 2000. A microsatellite linkage map of rainbow trout (*Oncorhynchus mykiss*) characterized by large sex-specific differences in recombination rates. *Genetics* 155: 1331–1345.
- Smith, P.** 2008. Antimicrobial resistance to antibiotics. *Rev. sci. tech. Off. int. Epiz.* 27 (1)
- Sommerset, I., Krossoy, B., Biering, E. & Frost, P.** 2005. Vaccines for fish in aquaculture. *Future Drugs, Expert Review of Vaccines* 4:89–101.
- Somorjai, I.M., Danzmann, R.G. & Ferguson, M.M.** 2003. Distribution of temperature tolerance QTL in Arctic charr (*Salvelinus alpinus*) and inferred homologies in rainbow trout (*Oncorhynchus mykiss*). *Genetics* 165: 1443–1456.
- Sønstebo, J.H., Borgstrøm, R. & Heun, M.** 2007. Genetic structure of brown trout (*Salmo trutta* L.) from the Hardangervidda mountain plateau (Norway) analyzed by microsatellite DNA: a basis for conservation guidelines. *Conservation Genetics* 8: 33–44.

- Starkey, W., Millar, R., Jenkins, M.E., Ireland, J.H., Muir, K. F. & Richards, R.H.** 2004. Detection of piscine nodavirus using real time nucleic acid based sequence amplification (NASBA) *Dis. Aquat. Org.* 59:93–100.
- Stokes, N.A. & Burreson, E.M.** 1995. A sensitive and specific DNA probe for the oyster pathogen *Haplosporidium nelsoni*. *J. Eukaryot. Microbiol.* 42: 350–357.
- Subasinghe, R.P.** 2009. Aquaculture development and blue revolution. *In Fisheries, Sustainability and Development*. Royal Swedish Academy of Agriculture and Forestry, pp. 281–301. Stockholm, Sweden.
- Subasinghe, R.P., Curry, D., McGladdery, S.E. & Bartley, D.** 2003. Recent Technological Innovations in Aquaculture. *Review of the state of world aquaculture*. FAO Fisheries Circular No. 886.
- Subasinghe, R., Soto, D. & Jia, J.** 2009. Global aquaculture and its role in sustainable development. *Reviews in Aquaculture* 1: 2–9.
- Tang, K.F.J. & Lightner, D.V.** 1999. A yellow head virus gene probe: application to in situ hybridization and determination of its nucleotide sequence. *Dis. Aquat. Org.* 35: 165–173.
- Tang, K.F.J., Pantoja, C.R., Poulos, B.T., Redman, R.M. & Lightner, D.V.** 2005. In situ hybridization demonstrates that *Litopenaeus vannamei*, *L. stylirostris* and *Penaeus monodon* are susceptible to experimental infections with infectious myonecrosis virus (IMNV). *Dis. Aquat. Org.* 63:261–265.
- Tenesa, A., Navarro, P., Hayes, B.J., Duffy, D.L., Clarke, G.M., Goddard, M.E., Visscher, P.M.** 2007. Recent human effective population size estimated from linkage disequilibrium. *Genome Research* 17:520–526.
- Thompson, K.D. & Adams, A.** 2004. Current Trends in Immunotherapy and Vaccine Development for Bacterial Diseases of Fish. *In* Leung Ka Yin, (ed). *Molecular Aspects of Fish and Marine Biology – Vol. 3 Current trends in the study of bacterial and viral fish and shrimp diseases*. Chapter 13.
- Thorgaard, G. H. & Gall, G.A.E.** 1979. Adult triploids in a rainbow trout family. *Genetics* 93: 961–973.
- Thorgaard, G. H.** 1986. Ploidy manipulation and performance. *Aquaculture* 57: 57–64.
- Uribe, P., & Espejo, R.T.** 2003. Effect of associated bacteria on the growth and toxicity of *Alexandrium catenella*. *Appl. Environ. Microbiol.* 69: 659–662.
- Uribe, P., Fuentes, D., Valdes, J., Shmaryahu, A., Zuniga, A., Valenzuela, P.D.T.** 2008. Preparation and analysis of an expressed sequence tag library from the toxic dinoflagellate *Alexandrium catenella*. *Mar. Biotechnol.* 10 (6): 692–700
- Valdebenito, I.** 2008. Hormone therapy for the artificial control of sexual maturity in fish culture: a review. *Arch. Med. Vet.* 40 N° 2, 115–123.
- Vasemägi, A., Nilsson, J. & Primmer, C.R.** 2005. Expressed sequence tag (EST)-linked microsatellites as a source of gene associated polymorphisms for detecting signatures of divergent selection in Atlantic salmon (*Salmo salar* L.). *Mol. Biol. & Evol.* 22: 1067–1076.
- Waldbieser, G.C., Bosworth, B.G., Nonneman, D.J. & Wolters, W.R.** 2001. A microsatellite-based genetic linkage map for channel catfish, *Ictalurus punctatus*. *Genet.* 158: 727–734.
- Wilson, K.J. & de la Vega, E.** 2005. The potential of microarrays to assist shrimp breeding and production: a review. *Australian Journal of Experimental Agriculture* 45: 901–911.

- Wilson, T. & Carson, J.** 2003. Development of sensitive, high-throughput one-tube RT-PCR-enzyme hybridisation assay to detect selected bacterial fish pathogens. *Dis. Aquatic Org.* 54: 127–134.
- Withler, R.E. & Beacham, T.D.** 1994. Genetic variation in body weight and flesh colour of the coho salmon (*Oncorhynchus kisutch*) in British Columbia. *Aquaculture*. 119(23): 135–148.
- Wright, J.M. & Bentzen, P.** 1994. Microsatellites – genetic markers for the future. *Rev. Fish Biol. & Fisheries* 4: 384–388.
- Young, W.P., Wheeler, P.A., Coryell, V.H., Keim, P. & Thorgaard, G.H.** 1998 A detailed linkage map of rainbow trout produced using doubled haploids. *Genet.* 148: 839–850.
- Zohar, Y. & Mylonas, C.C.** 2001. Endocrine manipulations of spawning in cultured fish: from hormones to genes. *Aquaculture* 197, 99–136.