



SporLaks – Industry-wide tracing of Norwegian farmed Atlantic salmon

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Atlantic salmon production in Norway

Every year 30 000 – 40 000 broodstock used to produce 350 million fish

Project Concept:

To use DNA parentage methods to trace escaped farmed salmon back to egg batch and/or cage of origin with 100% accuracy





Using DNA markers to trace fish in aquaculture



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www.elsevier.com/locate/aqua-online

Evaluation of three strategies using DNA markers for traceability in aquaculture species

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Simulations of the 'PAR' strategy

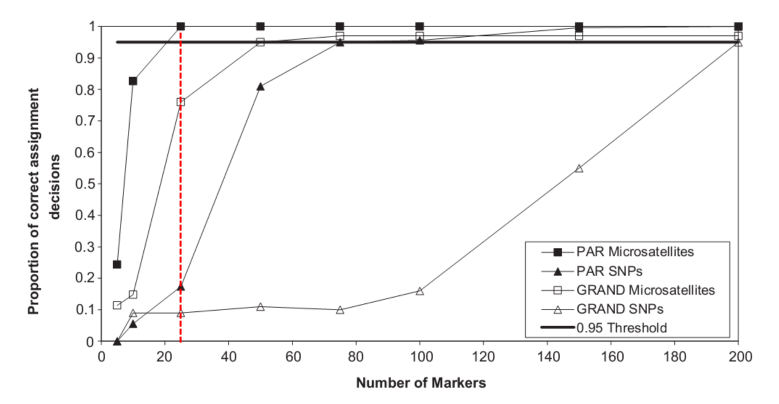
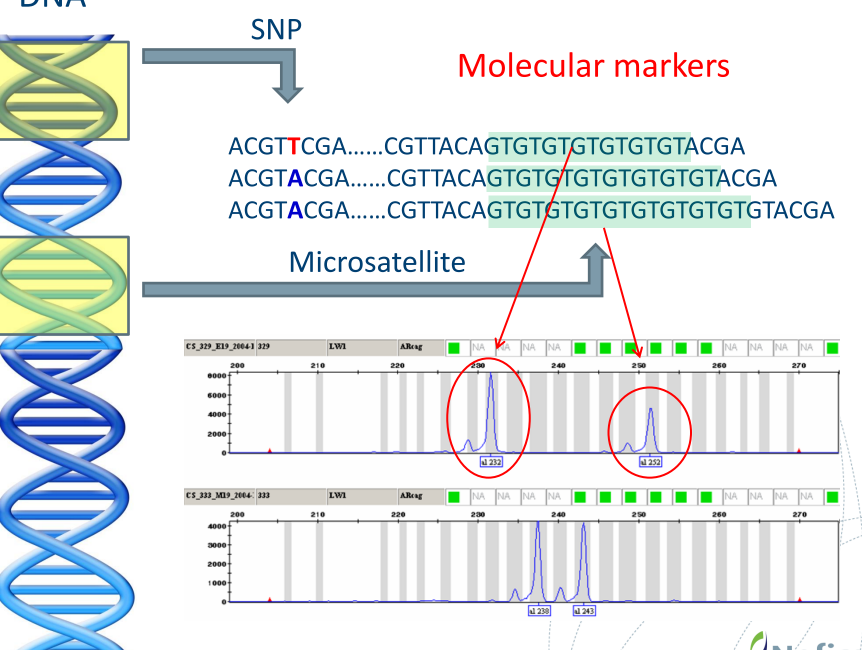


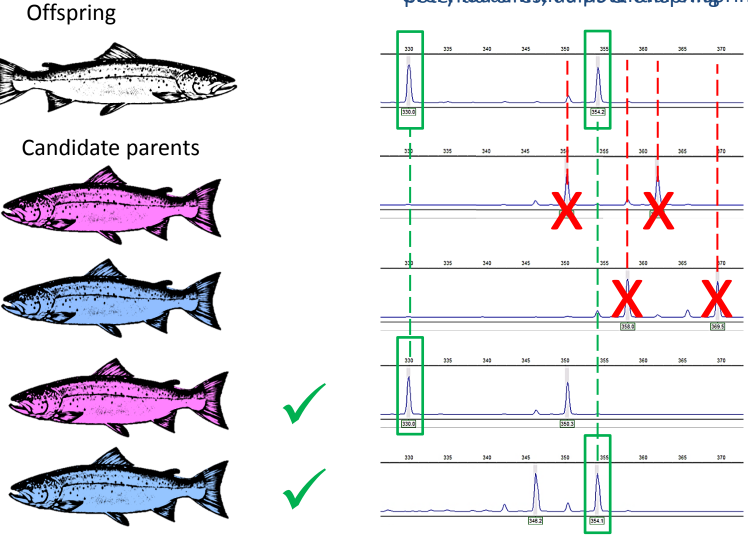
Fig. 5. Proportion of correct assignment decisions from strategies PAR and GRAND with increasing number of microsatellite and SNP markers.

DNA



Nofima

Parentage analysis – exclusion approach



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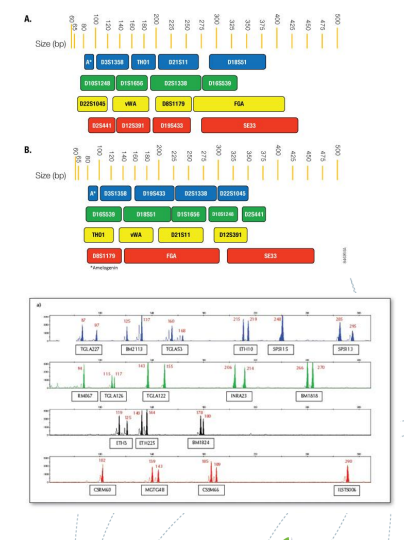
Multiplex PCR

- Amplify a 'panel' of markers together
- Very little DNA required
- Ability to handle mixtures and degraded samples
- Different fluorescent dyes used to distinguish alleles with overlapping size ranges

Established panels of microsatellites

- Human forensics/paternity
 - Promega PowerPlex

- Livestock and terrestrial species
 - Bovine Genotypes[™] Panel 3.1
 - Canine Genotypes[™] Panel 1.1
 - Equine Genotypes[™] Panel 1.1



Why don't we have such panels in aquaculture species?

- 1. Not so many top quality markers to choose from
- 2. Few concerted efforts to make highly optimised multiplexes

How to address this?

- 1. Use genome sequence data
 - No longer a shortage of markers to choose from
 - Tens to hundreds of thousands of microsatellites present
- 2. Use the same protocols/reagents as the optimised commercial panels



QDD pipeline using Atlantic salmon genome

BIOINFORMATICS APPLICATIONS NOTE

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Sequence analysis

QDD: a user-friendly program to select microsatellite markers and design primers from large sequencing projects

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80 markers

optimise multiplex primer design



Testing new markers with 'new generation' PCR reagents

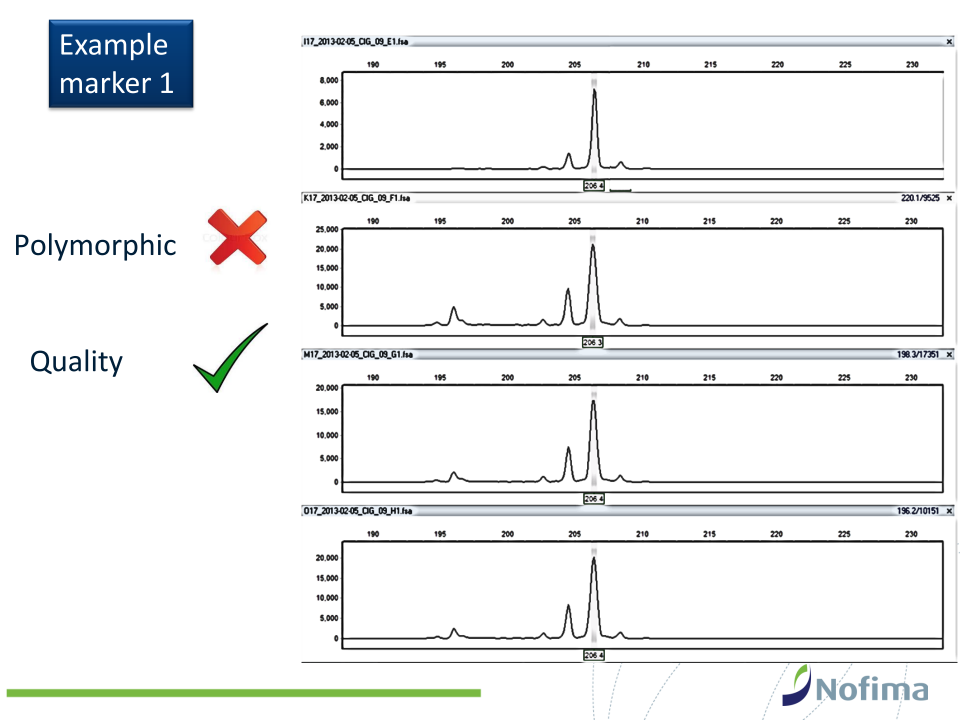
- Genotyping efficiency critical if thousands of samples are to be genotyped rapidly
- Commercial genotyping kits use 'advanced' reagents

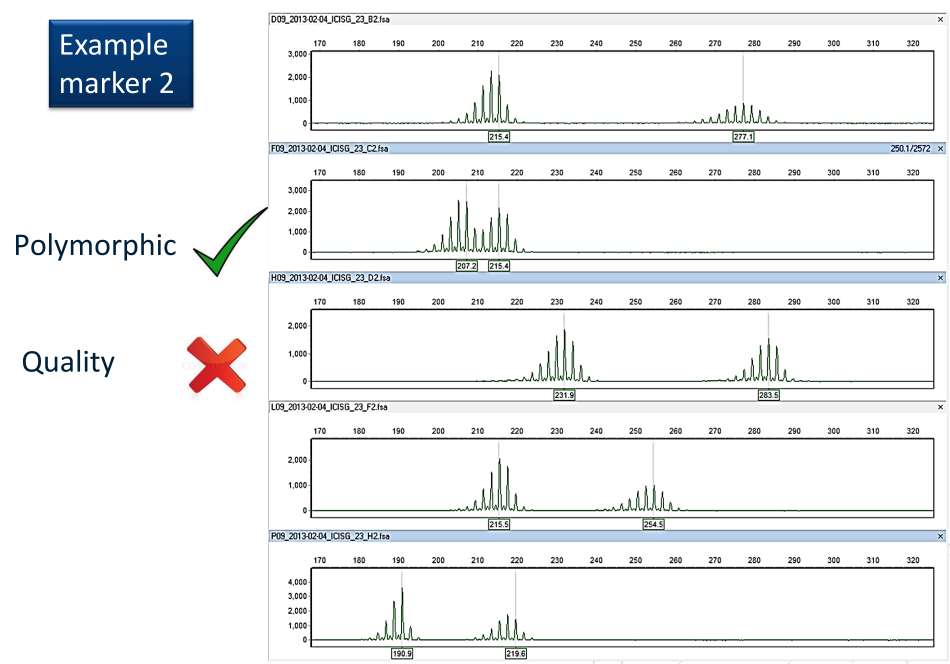
Q5 polymerase



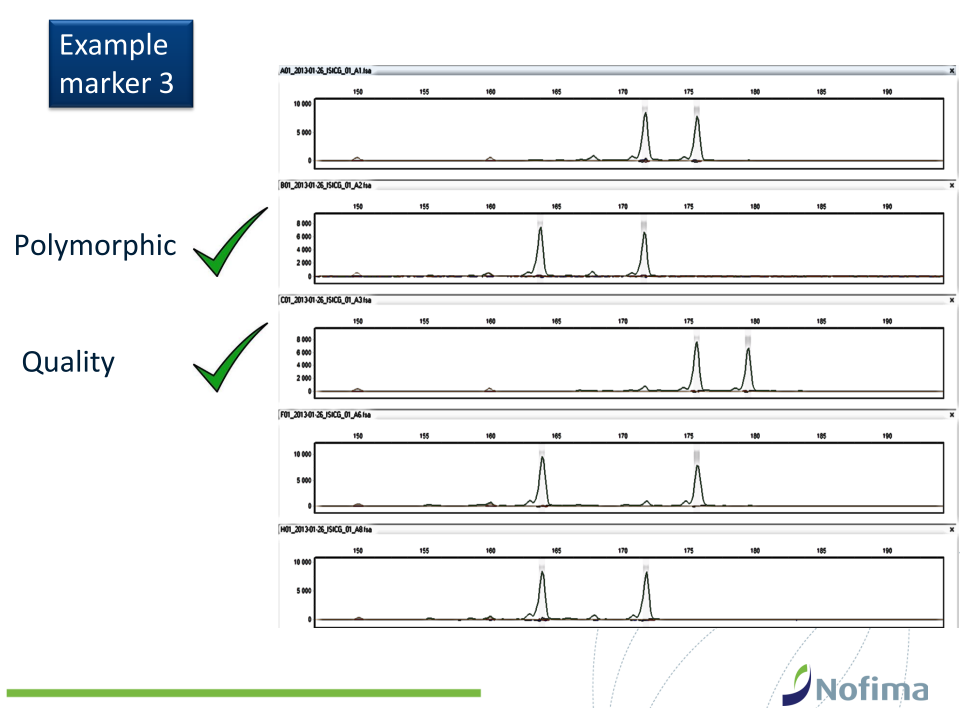
- Extremely high fidelity (>100X higher than Taq)
- Robust high specificity and yield with minimal optimization
- Very fast (10 s/kb)
- 30-40 minute two-step PCR program

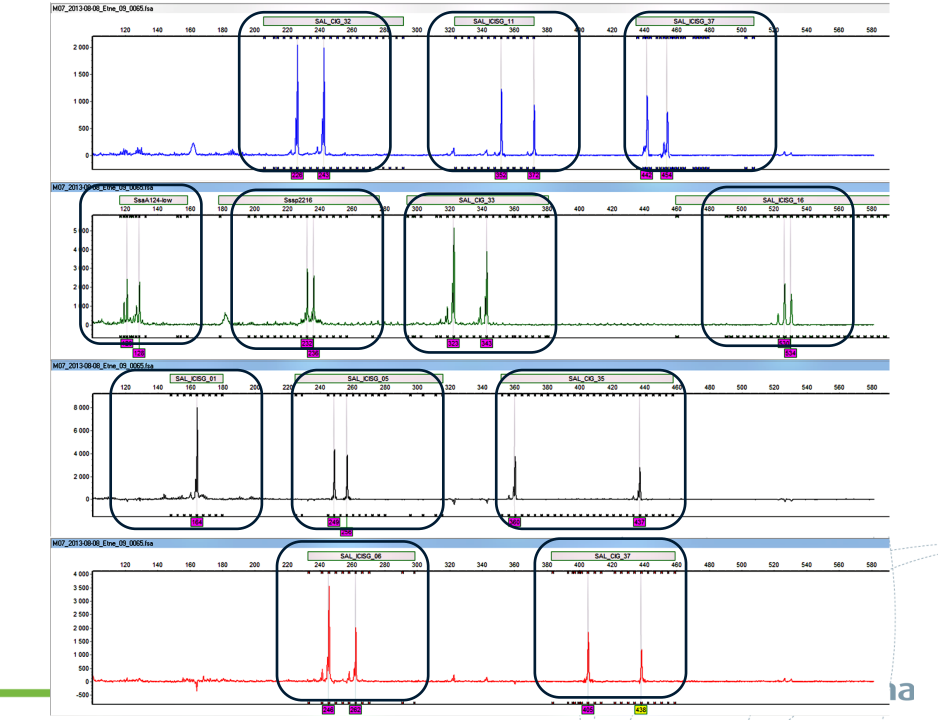


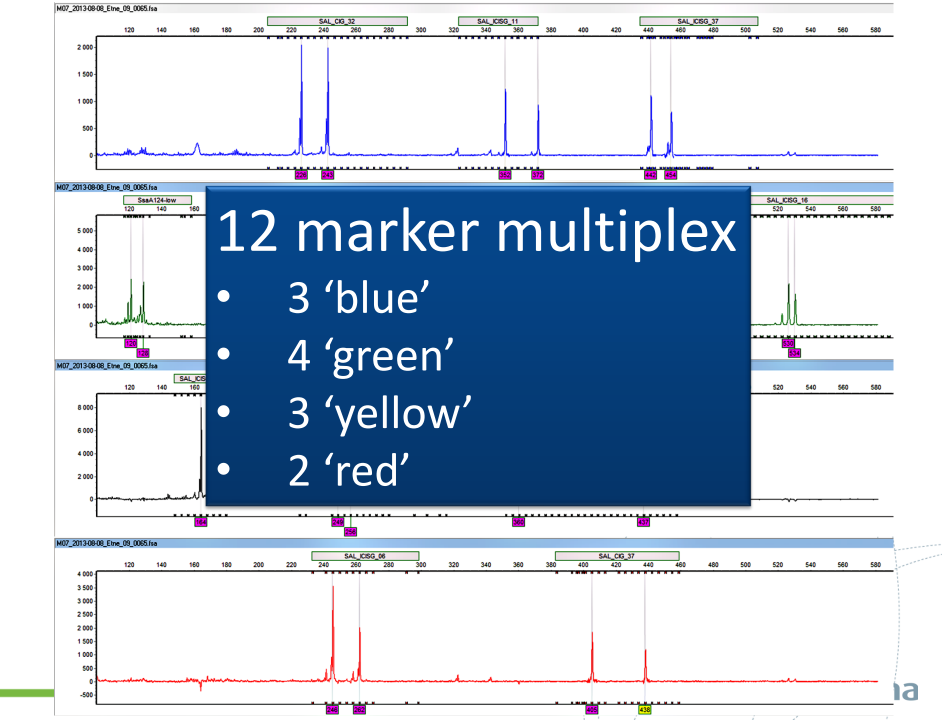








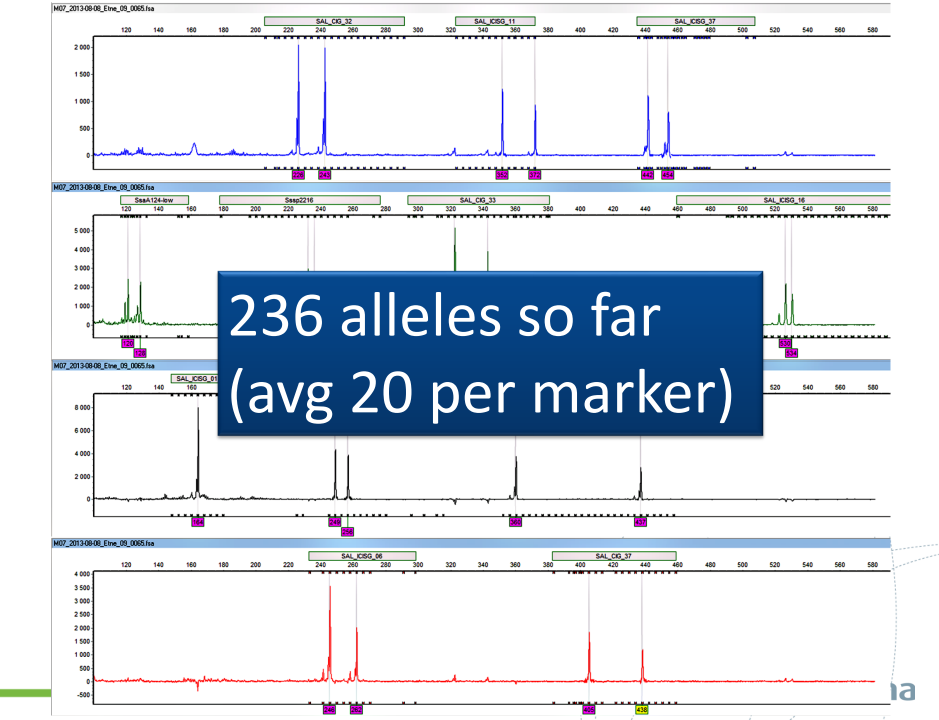




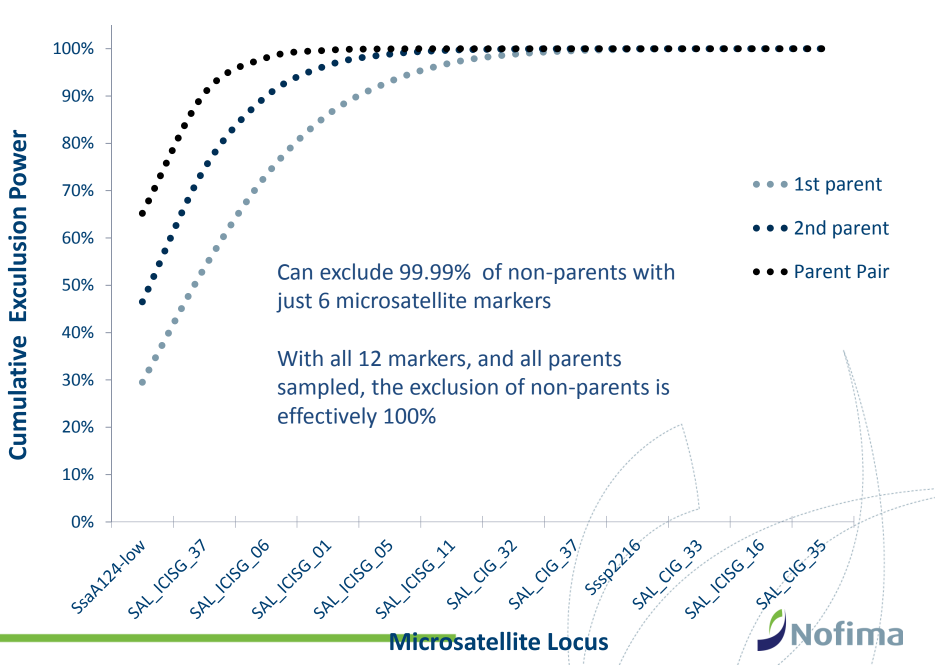
Wild fish sampling



- 17. Etne
- 18. Audna
- 19. Glomma



Parentage assignment accuracy



Validation study

 "Blind test" of parentage assignment using offspring from 230 Aqua Gen families (112 dams & 118 sires) + unrelated fish

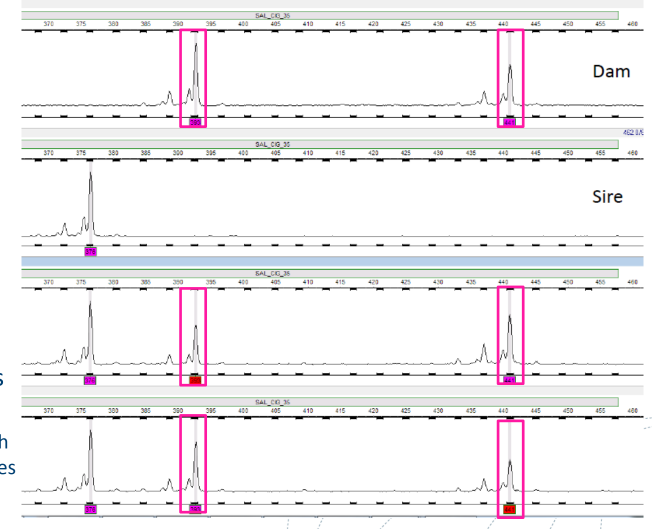
Validation set	Assigned to 1 or more parent	% Correct according to pedigree
520 offspring from 230 AquaGen families	519 (99.8%)	97%
40 unrelated AquaGen fish	0	100%
88 wild salmon	0	100%

- 1 dam couldn't be genotyped
- 504 offspring assigned to 2 parents, 15 assigned to a single parent
- 1 fish couldn't be assigned
- No un-related farmed fish & wild fish could be assigned to parents

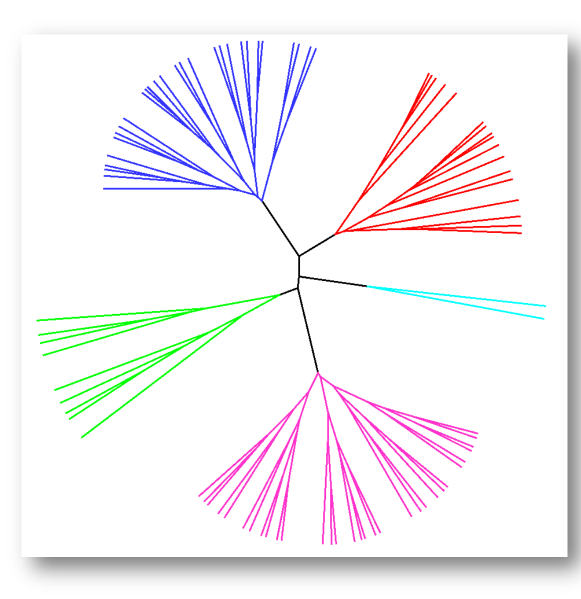


«Bonus» validation study results

- 4 of the wild fish had unusual genotype patterns
 - Further testing showed these were Salmon X Trout hybrids
- 2 of the Aqua Gen offspring had 3 alleles at most microsatellites
 - Assigned to parents
 - Offspring inherited both copies of mother's genes
 + 1 copy of father's



Relatedness clustering with multiplex



- Wild hatchery-reared smolt
- Progeny from single pair crosses of 5 males and 5 females
- Average relateness within each cluster (family) = 50%

High-throughput

Example for 384 samples

DNA extraction Crude method eg. Chelex

2 hours

PCR Q5 polymerase + robotics

Genotyping ABI 3730xl capillary sequencer



Logistical challenges

- Proposed tracing scheme will depend on sampling and genotyping of 50.000+ samples per year
- Huge logistical challenge
- Methods and protocols are needed to ensure:
 - Efficient sampling of thousands of fish by workers with a range of skill levels
 - Secure tracking, handling and transport of samples
 - Adequate preservation of tissue for downstream analysis
 - High throughput DNA extraction and genotyping

Sampling and sample storage

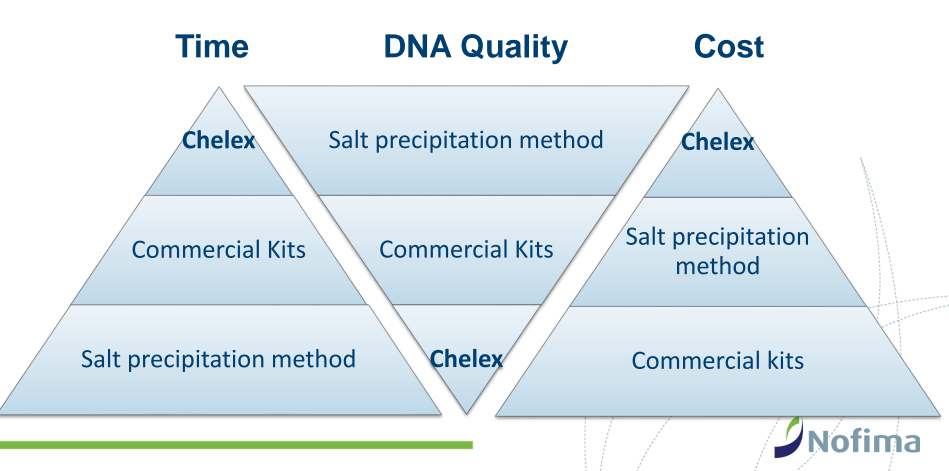
- Individual barcoded tubes in rack format
 - Read ID directly into database, no human error
- Room temperature storage in ethanol
- Compatible with lab robotics
- 'Biopsy' sampling ensures consistent sample size and good yield



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Genotyping effective with 'poor quality' DNA

- Tested three commonly used methods
 - Microsatellite genotyping performed well with Chelex[®]
 - cheapest, quickest, "roughest" method



Ongoing work

- Validate multiplex performance in different laboratories
- Marker data (allele number and frequencies) being fed into simulations
 - Test power of marker set at industry wide level





Conclusions

- Thousands of new markers identified from the Atlantic salmon genome
- Efficient multiplex of 12 high quality markers developed
- Protocol optimised for high-throughput genotyping at low cost
- Very high assignment power and very good assignment rates achieved for breeding companies
- Challenge lies in scaling up to whole-industry level and the logistics of sampling, genotyping, egg tracking
 - Rigorous database needed

