Recommendaions for reporting results of diagnostic genetic testing (biochemical, cytogenetic and molecular genetic)

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Genetic test results can have considerable importance for patients, their parents and more remote family members. Clinical therapy and surveillance, reproductive decisions and genetic diagnostics in family members, including prenatal diagnosis, are based on these results. The genetic test report should therefore provide a clear, concise, accurate, fully interpretative and authoritative answer to the clinical question. The need for harmonizing reporting practice of genetic tests has been recognised by the External Quality Assessment (EQA), providers and laboratories. The ESHG Genetic Services Quality Committee has produced reporting guidelines for the genetic disciplines (biochemical, cytogenetic and molecular genetic). These guidelines give assistance on report content, including the interpretation of results. Selected examples of genetic test reports for all three disciplines are provided in an annexe.

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Diagnostic genetic testing is an extremely rapidly expanding area encompassing a broad range of laboratory investigations to analyse chromosomes (from classical karyotype to molecular cytogenetics), nucleic acids (DNA, RNA), proteins and metabolites used to detect heritable or somatic mutations, genotypes or phenotypes related to disease and health. Genetic testing requires particular consideration in that it is usually performed only once in a patient’s lifetime, and the results may have considerable importance for lifetime decisions not only for the individuals being tested but also for children and family. Interpreting and reporting variation in germine chromosomes, DNA sequences or their products is a heavy clinical responsibility for prediction of susceptibility to disease, patient diagnosis, prognosis, counselling, treatment or family planning. Providing a set of reporting frameworks that can be customised for different testing contexts but share some common principles could be beneficial to the practice of a number of laboratories, including non-OECD members and/or laboratories that do not participate in External Quality Assessments (EQA), and to laboratories with blurred boundaries between research and genetic testing services.

Although several guidelines already exist for reporting the results of genetic testing,1,2,3 these focus on molecular genetic testing and do not cover the other two branches of laboratory genetics, namely biochemical genetics and cytogenetics. Based on recent surveys of EQA results presented by some European EQA providers and the request from genetic laboratories for comprehensive reporting guidelines, it was considered that a unifying attempt to harmonise the reporting practice of genetic tests in Europe and neighbouring countries would be welcome.

MATERIALS AND METHODS

On behalf of the Genetic Services Quality Committee of the European Society of Human Genetics (ESHG, https://www.eshg.org/), a workgroup composed of experts with long-term knowledge and experience in genetic testing was invited to several meetings, with the objective to answer the key question: is it feasible to classify a minimum set of issues and frameworks in reporting that would fit different testing scenarios encountered in cytogenetic, molecular and biochemical genetic laboratories? This workgroup included academic experts in clinical cytogenetics, molecular genetics, biochemical genetics, clinical genetics, bioinformatics, patient groups, accreditation bodies and EU organisations of EQA (proficiency testing) for genetic diagnostic laboratories. A preliminary draft including major key issues in reporting was discussed through meetings, then reviewed and updated by each of the participants, until reaching a consensus on shape and contents of these guidelines and recommendations.

Although these guidelines do not specifically address the reporting of results generated by massively parallel DNA sequencing, the reporting of the pathogenic variants identified by this method (or their absence) does not differ from reporting such variants found by conventional DNA sequencing.

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GENERAL COMMENTS
Scope of the document
These recommendations aim to provide guidance for reporting results of genetic testing by biochemical genetic, cytogenetic and molecular genetic techniques. They are distinct from (and bring additional information on reporting to) recently published documents such as good laboratory practices guidelines,4–10 quality assurance frameworks delivered by EuroGentest (www.eurogentest.org), evaluation frameworks and assessment of analytic validity developed by the Agency for Healthcare Research and Quality (AHRQ),11 CLIA (Clinical Laboratory Improvement Amendments) recommendations for good laboratory practices in molecular genetic testing12 or guidelines for licensing, certification or accreditation processes and general guidelines for quality assurance in molecular genetic testing provided by the Organisation for Economic Co-operation and Development (OECD).1 Reporting targeted genetic testing for monogenic disorders and for susceptibility traits with proven clinical utility, regardless of the extent of testing (that is, single gene or multiple genes) is within the scope of these recommendations.

Definition
Reports are specific formal medical documents from the laboratory to the referring physician and/or other health-care professionals regarding the results and interpretation of genetic testing in a patient and/or members of a family. The main goal of reports is to provide a clear, concise, accurate, fully interpretative and authoritative answer to a clinical question.

Minimal set of clinical information accompanying the samples
For the correct interpretation of genetic tests, a minimal set of clinical information from the referring physician or genetic counsellor must accompany the sample. The purpose of testing must be clearly identified (for example, confirmation of diagnosis that was established by other means, differential diagnosis of several disorders, carrier testing, presymptomatic testing and prenatal testing). The ethnicity of the individual tested should be included if relevant for the purpose of testing; this information may be useful to select the most appropriate test methods and to calculate residual risks. The referring physician should provide all clinical, laboratory, imaging, genealogic, genetic or other data that are relevant to the purpose of testing, and that may guide the laboratory in selecting the optimal laboratory procedures. For biochemical genetic testing, the referral should also include information on the conditions under which the samples were collected, on fed/fasting status and on the use of medications. For some clinical situations, it is essential to provide the laboratory with a family tree constructed by a genetic professional. For index cases, laboratories are strongly encouraged to request a more detailed disease-specific clinical form completed by the referring physician or genetic counsellor. Owing to potential deleterious consequences of shortcomings in the communication of critical information pertinent for the interpretation of results and clinical decision-making, some laboratories provide the ordering clinician with a clinical requisition form specific for the disease, which must be filled, signed and joined to the sample. Another benefit is to avoid non-medical office staff ordering genetic tests.13 If the purpose of testing or relevant clinical information is not provided, laboratories should request the missing clinical information required to perform the agreed testing.

Results of how many individuals in one report?
It is recommended that each unrelated patient should be reported in a separate and unique document, as many reports may ultimately be filed in individual patient or family files (for reasons of confidentiality). However, in recessive disorders it is appropriate (where legally permitted) to report the couple’s risk of having affected children together, which requires that both individuals are clearly identified.

When several family members are analysed simultaneously, policies vary as to whether they should be reported on the same or on different reports. This will depend on the disorder and the nature of the referral reason, as well as on the legislation on genetic testing in the respective country.

Guidelines for reporting
General guidelines and examples of reports are available in the Annex to these guidelines and on the ESHG, EuroGentest, CEQA (Cytogenetic European Quality Assessment), CF Network and ERNDIM (European Research Network for evaluation and improvement of screening, Diagnosis and treatment of Inherited disorders of Metabolism) websites for consultation.

CONTENTS OF THE REPORT
The following elements should be present in every report:

Administrative
Title (for example, ‘Result of molecular genetic analysis of the XX gene’ or ‘Results of enzyme assay for XX disease(s)’ or ‘Results of constitutional microarray analysis’).

The identity of the laboratory performing the analysis and issuing the report, with full contact details. If parts of analyses have been carried out in other laboratories, this fact must be clearly and unequivocally stated.

Full date of the report.
Page numbers indicating the total number of pages, essential when multiple pages are used. (for example, 1/1 or page 1 of 2).
Name and full address of the physician referring the patient.
Signature. The report must be electronically or manually signed by the authorised laboratory specialist who validated the analysis and interpreted the result; co-validation and co-signature by a second competent person is recommended (and mandatory in some countries). The name and function of signatories must be given.

Patient and sample identification
Patient identification:

- Full given name(s) and the surname (must be separate to avoid uncertainties, especially in sending samples to other countries).
- Unequivocal date of birth.
- Gender.

Fetal samples (CVS, amniocytes and so on) must be clearly distinguished from those of their mothers.

Note: Where legislation (for example, data protection legislation) prohibits the transmission of identifiable personal information, a code may be used instead of the patient’s name.

Ethnic/geographic origin, if relevant.

Date (and time of sample collection, if available; this information is crucial for biochemical genetic testing).

Date (and time) of sample arrival to the laboratory.

Information on the status of the sample if relevant (for example, frozen, decomposed, haemolyzed).

Material that has been tested (for example, ‘EDTA blood’, ‘cultured amniocytes’, ‘DNA extracted from…’, ‘heparin plasma’, ‘urine’ and so on). When the sample has been processed in another laboratory...
(for example, DNA has been extracted, plasma has been separated and frozen, amniocytes have been cultivated and so on) indicate precisely what was done with the samples and by whom (for example, sample: ‘DNA, extracted by XXXX, from blood’).

Unique sample ID number for each sample tested.

Patient identifiers should be included on each page of a multipage report.

Restatement of the clinical question
The interpretation of genetic test results depends usually on the clinical context. Therefore, in the section GENERAL COMMENTS above, a minimal set of clinical and laboratory data requested from the referring physician or genetic counsellor has been defined. Based on this information provided by the attending physician or genetic counsellor, the reports of genetic testing should explicitly restate the clinical question being asked. This usually comprises at least the following three elements:

The disease(s) or marker(s) being requested for analysis (for example, cystic fibrosis, lysosomal storage disorders, inborn errors of metabolism, chromosomal abnormalities);

The type of required testing (for example, diagnostic confirmation, carrier status, prenatal diagnosis);

The referral reason—why the request is being made (for example, positive family history, multiple congenital abnormalities, fasting hypoglycaemia and so on).

When the referral is due to abnormal results of previous genetic testing in either the same laboratory or a different laboratory, this fact must be clearly indicated in the report, with a reference to the relevant previous report.

Specification of genetic tests used
Gives useful, brief information on the methods used in the analysis. Additional information may be in footnotes.

Gives full details of the extent of the tests (for example, which exons screened for deletions in DMD, which FISH probes used, which mutations looked for in CF; resolution of cytogenetic analyses and so on). This information is particularly important when reporting normal/no abnormality detected results.

Where a commercially available kit is used, this should be clearly identified in the report, including the reference and version of the kit (and, where applicable, the list of mutations included in the version used).

For molecular genetic testing, gives the gene reference sequence that has been used to identify and describe the variants.

Where appropriate, gives the detection rate (sensitivity) in the population of origin of the individual tested.

Results
Present the results in a brief and unambiguous form. If several different tests have been performed, the results should be shown separately for each of the tests. The terms ‘positive’ and ‘negative’ can be ambiguous and should not be used.

For molecular genetic testing, reference sequence (including Build No.) and nomenclature of detected genetic variants should be meaningful, unambiguous and consistent using the recommended HGVS nomenclature valid at the time of reporting. As both HGVS nomenclature and the reference sequences change in time, other descriptions of the variant that have been widely used in the past may be given in parentheses, if appropriate.

If more than one possibly pathogenic variant has been detected, a comment on the phase of variants (which may be unknown) should be added, taking into account the expected mode of inheritance of the disorder.

For cytogenetic and microarray testing, the most recent International System for Human Cytogenetic Nomenclature (ISCN) nomenclature should be used.

It may be useful to give a list of genes involved (by HGNC symbol and/or OMIM number) for abnormal microarray results and for other tests involving multiple genes.

For biochemical genetic testing using qualitative tests, the report should clearly distinguish normal results from nonspecific findings and from clearly abnormal results (including an indication of which diseases are most likely). For quantitative analyses of single analytes, both the value and the reference range should be given. For enzyme assays it is usual to show, in addition, the activities of controls run in parallel with the patient sample; reports on enzyme activity should contain the E.C. number of the enzyme. For quantitative profile analyses, the report should only summarise the major findings distinguishing normal, nonspecific and disease-specific profiles; numeric values of the profile analyses should be sent as attachments.

Interpretation of results
General recommendations for the interpretation of genetic laboratory results are given in this section. The Quality Committee realises that differences between countries may exist, as to where the boundary of the responsibilities of laboratory specialist versus the clinician lies. For example, the use of prediction programs to predict the effect of a novel missense variant or a variant that may affect splicing is evidently the expertise and responsibility of the laboratory specialist. The same holds true for the interpretation of intermediate biochemical results, although the clinical phenotype may sometimes be decisive for the interpretation. On the other hand, relating the phenotype to what is known in literature on the effect of a rare copy number variant will highly depend on the background of the laboratory specialist (in some countries, cytogeneticists are clinical geneticists) and the referring clinician. With the use of new high-resolution and high-throughput sequencing techniques, the number of (copy number) variants of unknown clinical significance will increase, and the prediction programs and the databases with normal variants have their limitations. Thus, laboratory and clinical specialists should discuss and document what the minimal interpretation in the laboratory report should be. In the opinion of the Quality Committee, the report should at least contain all relevant information that enables the clinician to perform the clinical interpretation using literature resources (for example, which RefSeq genes are located in a copy number loss, or what the results are of prediction programs on a missense variant and whether or not it has been reported in databases like dbSNP). It may be worthwhile to inform the clinician that genealogical/family studies may be helpful in the interpretation, and that the interpretation is based on current knowledge and thus may change in the future due to additional evidence.

Although this is beyond the scope of these recommendations for reporting results of diagnostic testing, the committee acknowledges the need for curated locus/disease-specific databases and databases of copy number variants. Such databases should be supported by an international expert committee (as proposed by Tavtigian et al.16).

The report must provide a full and clear interpretation of results, depending both on the clinical context and the reason for referral (for example, diagnostic test, carrier test, prenatal test). Reports on patients may be read by a variety of professionals involved in their care, many
of whom will be unable to interpret fully the results of genetic testing. In order to provide a full interpretation, the results must be viewed in the context of the relevant clinical and family information available (for example, relationship between the patient and the index case where relevant, family pedigree, ethnic background, clinical and laboratory data provided by the referring physician and others).

If more tests have been carried out using the same sample(s), the interpretation part of the report should integrate the individual results and should answer the clinical question in a clear and concise way. This integration of findings with relevance to the tested individual is especially important when multiple susceptibility genetic variants have been analysed.

Answer the question in a concise and clear way, after taking into account any appropriate additional information supplied. It is recommended that laboratories highlight this conclusion (bold, underline, large font, text in a box and so on). In particular, remember that negative results (‘no mutation detected’) can easily be misinterpreted by non-specialists as ‘exclusions’. Significance of results in respect to the referral reason (clinical question) should be clearly stated and should belong to one the following five categories.

In some cases there may be more than one conclusion:

- **Normal finding(s) (physiological finding, normal variation):** findings within the range of physiological variation for the given individual considering ethnicity, age, sex, maturity and other relevant factors; for example, nucleic acid sequence corresponding to the reference sequence or containing genetic variants considered usual variations at the time of reporting, chromosomal heteromorphisms as defined in the ISCN, frequently occurring CNVs reported in the database for genomic variants (DGV), concentrations of analytes in quantitative biochemical genetic tests within reference ranges, results of qualitative biochemical genetic tests showing a usual result/profile.

  If relevant, it may be important to provide an estimate of the diagnostic sensitivity (that is, the proportion of affected individuals likely to be detected; or one minus the false-negative rate). It can be useful to provide a key reference to support sensitivity estimates, when appropriate, but the report should not become a scientific discussion.

- **Non-specific finding(s) without clinical relevance:** findings outside of the physiological variation but not associated with a known disease; for example, elevation of multiple serum amino acids in a patient receiving parenteral nutrition, presence of drugs and their metabolites in biochemical tests, signs of sample decomposition. Such findings should only be reported if they may be relevant to the result (for example, issues of sample quality).

- **Incidental finding(s) with possible clinical relevance:** findings indicating a clinically relevant issue, but unrelated to the clinical question that was asked (for example, signs of sex chromosome abnormality when analysing an X-linked disorder, evidence of predisposition to an unrelated condition). The decision on whether to report such findings will depend on local policy and on how the patient has been counselled about this possibility. A clear policy on reporting incidental findings should be in place in all institutions offering genetic testing.

- **Finding(s) of uncertain significance:** findings outside of the physiological variation but with possible or putative relevance to the clinical question asked; for example, novel missense variants, novel putative splicing variants with unproven effect at RNA level, mosaicism or novel rearrangements in cytogenetic analyses, copy number variations and intermediate analyte concentration/enzyme activity, suggesting either heterozygosity or a mild phenotype. It is important that findings of uncertain significance are included in reports, as their significance may become clear at a later date.

Attempts should be made to explain significance of the finding using literature, additional databases, prediction programs and other means, and to report the likelihood of the finding in respect to the clinical question asked. In some cases, it may be important to provide an estimate of diagnostic specificity, which indicates the risk of a false-positive result (for example, in late-onset diseases or in cases of reduced penetrance). When practical and appropriate, the advice of a Clinical Geneticist should be sought.

- **Pathognomonic (disease-specific, pathological) finding(s):** findings that are outside of the physiological variation and that are, at the present state of knowledge, unequivocally associated with a clinically relevant disease or group of diseases. Pathogenicity of findings should be clearly supported from data in literature/databases, segregation analysis, functional tests of genetic variants and other relevant means.

  Description of genotype can include the correct assignment of phase, which may require the study of familial segregation of alleles. When appropriate, reports must explicitly mention that ‘both parents should be tested to confirm their carrier status and to provide formal confirmation of the diagnosis of their child.’

  When appropriate or required by the requesting physician, genetic carrier risks should be stated. Risk estimates may require the application of Bayesian calculations.

  It should be indicated if further tests could be undertaken to improve the accuracy or scope of the interpretation. This may include tests for additional disorders or additional tests to more fully investigate the disease in question. If the additional tests suggested are not performed ‘in-house’, alternative specialist laboratories where the sample may be sent may be proposed.

  If any other information could be supplied or obtained by the referring clinician that might improve the accuracy of your interpretation, it should be stated in the report (for example, arranging testing of the index case in a family to confirm a diagnosis or to determine which mutations are present).

  References: It may be useful to quote appropriate references for the interpretation of results, particularly of rare pathogenic and unclassified variants, as well as for certain risk calculations.

  **Requirement for genetic counselling:** The report should carry the reminder to the referring physician that ‘genetic tests should be accompanied by genetic counselling’ (or similar). In certain cases (prenatal diagnosis, presymptomatic testing or results with major implications for other family members), it is appropriate to indicate that ‘this result and its implications should be transmitted only in a specialized genetic consultation’ (or similar).

  **Family follow-up:** When a new diagnosis is made (positive result in a diagnostic confirmation), it is appropriate to state specifically that the result has ‘potentially important implications for other family members’ or equivalent. Depending on the context, it may be appropriate to explicitly mention the recommendation to test the partner, the possibility of cascade screening tests in relatives, the possibility of prenatal diagnosis or preimplantation genetic diagnosis. Where appropriate, the risk for future offspring should be calculated and provided (using carrier frequencies from an appropriate population).

  It should never be stated that prenatal or presymptomatic diagnosis is ‘indicated’ or ‘necessary’. The role of the laboratory does not go beyond stating that prenatal/presymptomatic diagnosis would be possible.

**INTERIM REPORTS**

It may in some circumstances be useful to issue a report before all studies are complete (for example, when indicative preliminary results
have been obtained, but a long delay is expected before the final results will be ready). Interim reports should be clearly marked as such and should be worded to avoid misinterpretation of their status. Thus, phrases or summary statements appearing to give a definitive result should not be used. It should be clearly stated which analyses are still underway.

The definitive report should clearly state which are the new results, and should include a general conclusion, taking account of all results.}

**REPORTING MULTIPLE PATIENTS**

As a general rule, each *unrelated* patient should be reported on a separate and unique document, as the reports will ultimately be filed in individual patient or family files (as well as for reasons of confidentiality).

When several family members are analysed simultaneously, policies vary as to whether they should be reported on the same or on different reports. This will depend on the disorder, on the nature of the analysis and also on the legislation (in some countries, familial analysis and prenatal diagnosis for hereditary disorders can be referred only by clinical geneticists, and family files are stored only in genetic centres, and only relevant information is delivered to each individual).

**Predictive test results** must always be reported as separate, individual reports.

**Linked-marker studies** are only useful in the context of alleles inherited by several family members, which must be included in the report. It is recommended that the number of individuals reported should be limited to only those essential for accurate analysis. An interpretation and final risk should be given for only one person per report.

For *prenatal diagnosis*, it is recommended that the report includes only the result of the fetus. Parental results should be cross-referenced, but their results reported separately.

In *carrier testing for a recessive disorder* for a couple, the test results for one partner must be interpreted in light of the other partner’s results. Laboratories may issue a single report, or separate reports with cross references to the partner’s results (as the couple may separate). It is recognised that, in some jurisdictions, it is not permissible to mention more than one individual in a clinical report.

Where the result of the parents is needed to interpret the result in the child, only the array result of the specific CNV(s) in the parents is given in the child’s report, and not the full array results.

**DISCLAIMER**

Disclaimers should only be included where they are relevant and useful; inclusion of multiple disclaimers on every report whether relevant or not is discouraged.

Where relevant, mention the **possibility of errors** due to factors beyond the control of the laboratory (for example, the risk of ‘non-paternity’ and the need for family relationships as stated on the referral forms being correct; limited validity of biochemical testing if pre-analytical conditions were not well controlled).

In indirect (linkage) analyses, it is sometimes advisable to state that the ‘accuracy of the result depends on the clinical diagnosis and the assumption that gene X is responsible for the disease’ (or similar). Laboratories might wish to add a note of caution when reports are based on DNA samples or reports sent from another laboratory.

It may be advisable to add a standard phrase, indicating that ‘this report may not be copied or reproduced, except in its total’.

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Annexes

1. Selected examples of reports

Unrelated examples of typical report elements are shown below. Additional report examples may be available from some EQA providers.

A. Biochemical genetic testing

Example 1.

Results of organic acid analysis in urine.

Laboratory of Inherited Metabolic Diseases, City, Country
Contact: Prof. xxx, telephone xxx-xxxx-xxxx
March 2, 2013
Referring physician: Dr.X, Hospital, Street, City, Country

Patient and sample information

Patient name: John DOE D.O.B.: March 2, 2011
Gender: Male Laboratory ID number: U235/13
Sample collected on: February 28, 2013
Sample arrival at laboratory: March 1, 2013, 10:15
Sample type: early morning urine

Referral reasons

Unexplained severe hypoketotic hypoglycemia upon prolonged fasting. Suspected fatty acid oxidation defect or other inborn error of metabolism.

Methods:
Analysis of the profile of organic acids in urine (ethyl acetate extraction, TMS derivatization with oximation, GC-MS analysis).  

Results:
The analysis of organic acids in urine reveals moderate dicarboxylic aciduria (adipic, suberic and sebacic acids) with traces of hexanoylglycine and suberylglycine, and with no increase of glutarate. See attachment for list of analytes, their concentrations and reference ranges.

Interpretation:
The profile of organic acids in urine is consistent with medium chain acyl-CoA dehydrogenase deficiency (OMIM 201450). Diagnosis should be confirmed by analysis of acylcarnitines in blood and/or molecular genetic analysis of the ACADM gene.

Signed

Head of Lab
Example 2.

Results of cystathionine beta-synthase assay in cultured fibroblasts (E.C. 4.2.1.22)

Laboratory of Inherited Metabolic Diseases, City, Country
Contact: Prof. xxx, telephone xxx-xxx-xxxxxx
February 2, 2013
Referring physician: Dr.X, Hospital, Street, City, Country

Patient and sample information
Patient name: Joan HAWKES
D.O.B.: April 12, 2009
Gender: Female
Laboratory ID number: F2/13
Sample collected on: January 13, 2013
Sample arrival at laboratory: January 13, 2013
Sample type: cultured skin fibroblasts obtained from tissue culture laboratory of Pediatric University Teaching Hospital, Churchville (ID number UTC345/2012)

Referral reasons
A patient with developmental delay and lens subluxation with severe hyperhomocysteinemia with hypermethioninemia. Suspected cystathionine beta-synthase deficiency.

Methods:
Activity of CBS was determined in cell-free extract of subcultured skin fibroblasts. The assay is based on the conversion of $[{}^{14}C]$-serine and homocysteine to labelled cystathionine and chromatographic separation of product.\textsuperscript{16}

Results:

|                      | CBS activity (nmol cystathionine/mg protein/hour) |
|----------------------|--------------------------------------------------|
|                      | -pyridoxal-phosphate                              |
| Patient              | 0.12                                              |
| Experimental control | 8.6                                               |
| (in series)          | 10.3                                              |
| Reference range      | 4.1-21.6                                          |
|                      | 5.3-24.4                                          |
| (n=35)               |                                                   |

The values are averages of those obtained in two batches of cells assayed in duplicate.

Interpretation:
The CBS activity in cultured fibroblasts is markedly decreased in the range typical for homozygous deficiency. This result confirms homocystinuria due to CBS deficiency (OMIM 236200). To facilitate genetic counselling in the family molecular genetic analysis of the CBS gene can be performed.

Signed

Head of Lab
B. Cytogenetic testing

Example 1.

| Logo                                      | Laboratory header | Accreditation logo and number |
|-------------------------------------------|-------------------|-------------------------------|
| Referring clinician :Doctor X             |                   | Laboratory XXX               |
| Street, City, Country                    |                   | Department of Cytogenetics   |
|                                           |                   | Address                       |
|                                           |                   | Contact person                |
|                                           |                   | Telephone – Fax               |

Name: John SMITH
Gender: Male
Nature of sample: DNA (extracted from blood at xxx, UK).
Date of reception of sample: 18/02/2013

Date of Birth: 07/02/2002
Ethnic origin: Caucasian
Internal ID N°: M1253/12
Date of report: 18/03/2013

Cytogenetic Analysis

Referral reason
10 year old boy with multiple congenital abnormalities including x, y and z. Microarray analysis requested.

Result:
arr[hg19] 3q25.2q26.31(153,508,217-173,161,913)x3.

Array CGH analysis has shown a male hybridisation pattern with an approximately 19.6Mb gain of chromosome 3, bands q25.2-q26.31.

Interpretation:
- This region contains xx OMIM listed genes and given its size and gene density this 3q25.2q26.31 gain is the likely cause of John’s clinical phenotype. *(List the appropriate genes).*
- Patients with 3q duplications overlapping this case have been reported in xx and xx databases (or reference) with clinical features of u,v,x,y, and z.
- The 3q gain may have arisen as a new mutation or inherited from a parent who may carry a balanced rearrangement. In order to investigate the inheritance of this gain and inform recurrence risk, parental blood samples (5ml in Lithium heparin) are requested for cytogenetic studies in addition to a repeat (Lithium heparin) blood sample for John.
- John is at 50% risk of transmitting this gain to his children and this should be discussed at a suitable age, if considered appropriate.
- You may wish to refer this patient and his parents to the Clinical Geneticist to discuss the result and recurrence risk.
- Array data will be uploaded anonymously into the xx database. Please find enclosed a patient information and consent form which needs to be completed for data to be shared with other xx consortium members. This may help in further interpretation of this case.

**Analysis by:** <name and signature>  
**Authorised by:** <name and signature>  
**Designation (e.g. Biologist or Section Head)**  
**Designation (e.g. Biologist or Director)**

Method: Microarray analysis using Oligo platform (version, manufacturer), build no. Aberrations below 150kb are not reported.

Limitations of test: This technique will not detect low level mosaicism, balanced translocations and point mutations.

The analysis also identified several copy number variations (CNVs), which were all described as variants in the Database of Genomic Variants (http://projects.tcag.ca/cgi-bin/variation/gebrowse/hg19/). A list of these CNVs is available upon request.

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**Example 2.**

| Logo | Lab Header | Accreditation logo and number |
|------|------------|------------------------------|
| Referring clinician: Doctor X | Laboratory XXX | Department of Cytogenetics |
| Street, City, Country | Address | Contact person |
| | Telephone – Fax | |

**Patient identification**

Name: Mary SMITH  
Gender: Female  
Nature of sample: Blood  
Date of reception of sample: 18/02/2013  
Date of Birth: 07/01/1983  
Ethnic origin: Caucasian  
Internal ID N°: T1200/12  
Date of report: 18/03/2013

**Cytogenetic Analysis**

**Reasons for referral**

Recurrent miscarriages (x3). Products of conception from 16 week fetus showed a der(4)t(4;12)(p14;p13.3).

**Result:**

46.XX,t(4;12)(p14;p13.3).ish t(4;12)(WHSC1-;WHSC1+).

Chromosome analysis revealed a female karyotype with a balanced translocation with the breakpoints in the short arm of chromosomes 4 (p14) and the short arm of chromosome 12 (p13.3).

The 4p translocation breakpoint was proximal to the Wolf-Hirschhorn syndrome locus WHSC1:

**Interpretation:**

- The presence of the translocation in Mary would explain the recurrent miscarriages of the couple.
- Translocations in general have an increased risk for embryos with an unbalanced karyotype, depending on the segregation behaviour of the chromosomes involved. This chromosome result has an increased risk for spontaneous pregnancy losses or for live-born children with an unbalanced karyotype, which may result in developmental delay and congenital anomalies. The risk for each of these outcomes in this couple can be estimated by a clinical geneticist after full information of the family pedigree has been obtained.
- Mrs. Smith can opt for prenatal diagnosis in future pregnancies
- Healthy children of this couple may also carry the balanced translocation and chromosome analysis can be offered to them as soon as they are of reproductive age.
- The translocation t(4;12) could have arisen de novo in Mary but may also be present in one of her parents and thus also in her siblings and other relatives. Family studies can be initiated by the clinical geneticist/genetic counsellor.

**Analysis by:** <name and signature>  
**Authorised by:** <name and signature>  
**Designation (e.g. Biologist or Section Head)**  
**Designation (e.g. Biologist or Director)**  

**Methods:** Chromosomal analysis using G-Banding and FISH

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C. Molecular genetic testing

Example 1.

| Recommendation for reporting results of diagnostic genetic testing |
|---------------------------------------------------------------|
| **Name:** William LEWIS |
| **Gender:** Male |
| **Sample Type:** EDTA Blood |
| **Sample Date:** 20/02/2013 |

**MOLECULAR GENETIC ANALYSIS FOR CYSTIC FIBROSIS (CFTR gene)**

**REFERRAL REASON:**

CF carrier status and risk of having a child with CF. William’s sibling died of CF; William is in good health. No molecular studies have been done in the family. Based on this information, William has a prior risk of 2 in 3 of carrying one of the familial CF mutations. William’s wife Florence Lewis (d.o.b.12/17/1999) who is French, has no family history of CF and is in good health.

| RESULT |
|---|
| c.[1585-1G>A];[=] |

(former nomenclature: 1717-1G>A/n)

| SUMMARY |
|---|
| William LEWIS carries the CF mutation c.1585-1G>A; partner testing recommended |

*Key: [=] and n denote normal allele present*

**REPORT INTERPRETATION:**

- This individual is heterozygous for the c.1585-1G>A CF mutation. The remaining mutations listed below are absent and together, these mutations account for approximately 93.5% of the CF mutations seen in the local population.
- The risk for this couple of having a child with CF, based on the result of molecular analysis in William LEWIS, is around 1/130 (1/2x1/2x1/32-a priori CF carrier frequency in individuals from France).
- The risk for future offspring of this couple could be determined more accurately if his wife could be tested. It is recommended that testing for the most common CF mutations in the French population should be offered to Florence to determine their combined risk as a couple of having a child with CF.
- All of this individual’s children have a 1 in 2 risk of inheriting the CF mutation c.1585-1G>A. Carrier testing for the c.1585-1G>A mutation for this individual’s children would be available once they reach adulthood.
- Carrier testing for the detected mutation could be offered to the adult relatives of this individual based on this information.
- As this result has important implications for this individual and their family, a referral to a clinical genetics service is recommended.

| REPORTED by: |
|---|
| <Name and signature> |

**Method:** Analysis based on allele-specific oligonucleotide hybridisation using the NNN CF analysis platform v2.1. Mutations analysed: p.Ala455Glu (A455E), p.Ala559Thr (A559T), p.Phe508del (ΔF508), p.Ile507del (ΔI507), p.Gly858Glu (G858E), p.Gly551Asp (G551D), p.Gly542X (G542X), p.Met1101Lys (M1101K), p.Glu1303Lys (N1303K), p.[Arg117His] (R117H), p.Arg334Trp (R334W), p.Arg477His (R477H), p.Arg477Pro (R477P), p.Arg553X (R553X), p.Arg560Thr (R560T), p.Arg1162X (R1162X), p.Ser549Asn (S549N), p.Ser549Arg (S549R), p.Ser1255X (S1255X), p.Val520Pro (V520F), p.Trp1282X (W1282X), p.Tyr122X (Y122X), p.Tyr1092X (Y1092X), p.Leu88fs (394delTT), c.489+1G>T (621+1G→T), c.579+1G>T (711+1G→T), p.Phe316fs (1078delT), c.1585-1G>A (1717-1G>A), c.1766+1G>A (1898+1G>A), c.1766+3G>T (1898+5G>T), c.2051+2052delAAnG (2183AA>GG), p.Leu684fs (2184delA), p.Glu726fs (2307insA), p.Glu726fs (2307insA), c.2657+5G>A (2781+5G>A), p.Arg117fs (3659delC), c.3717+10kbC>T (3849+10kbC→T), p.Leu1258fs (3876delA), p.Leu1258fs (3905insT).

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Together these mutations account for 93.5% of mutations seen in the local population.

The mutations are named using CFTR sequence accession number NM_000492.3 (gi:90421312). The current description of the mutations are as recommended by HGVS, (www.hgvs.org/mutnomen)

Please note that DNA has been stored from this patient’s sample at this centre, and will be kept indefinitely unless a written request for its disposal is received from the patient or their parent/guardian. Individual elements of this report should not be copied or transferred to other systems; the report should only be copied in its entirety.

Filename: 12cf7677

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**Example 2.**

| Logo | Lab Header | Accreditation logo |
|------|------------|--------------------|
| Referring clinician: Doctor X | Laboratory XXX | |
| Clinic, Street, City | Address | |
| | Contact person | |
| | Telephone – Fax | |

**Name:** Nancy DAUTARD  
**Gender:** Female  
**Sample Type:** EDTA Blood  
**Sample Date:** 20/02/2013

**Date of Birth:** 05/06/1978  
**Pedigree No:** DMD 742  
**Laboratory Number:** D 03988  
**Date of report:** 18 March 2013

### MOLECULAR GENETIC ANALYSIS of the DMD GENE

**REFERRAL REASON:**
Carrier testing for Duchenne Muscular Dystrophy (DMD). Nancy DAUTARD is the mother of Jérôme DAUTARD (DOB 06/05/2008) who has been recently reported by our laboratory with a hemizygous duplication of exons 2 to 4 of the dystrophin gene (DMD gene). Nancy DAUTARD has normal CK levels.

| RESULT |  |
|--------|---|
| - Abnormal dosage of exons 2 to 4 in the DMD gene, indicating a heterozygous duplication of these exons in the DNA extracted from the leucocytes of Mrs Nancy DAUTARD. |
| - Normal amplification for exons 1 and 5 and for internal controls. |

| SUMMARY |  |
|---------|---|
| Nancy DAUTARD carries the duplication of exons 2-4. |

### REPORT INTERPRETATION:

- Mrs Nancy DAUTARD (DOB 05/06/1978) carries a heterozygous duplication from exon 2 to exon 4 in the DMD gene.
- **She is a carrier of DMD. She is at risk of having further affected sons (50% risk) or carrier daughters (50% risk).**
- Genetic counselling is recommended.
- Direct prenatal diagnosis can be performed in future pregnancies.
- Direct carrier testing can be offered to female relatives following referral to a clinical genetics service.

**Reported by:** <Name and signature>  
**Checked by:** <Name and signature>  
**Molecular Geneticist**  
**Director or Second Molecular Geneticist**

**Method:** DNA extracted using the Flexpex kit from Qiagen.  
**Screening method:** MLPA (Multiplex Ligation-dependent Probe Amplification/MRC Holland, SALSA kit PO34-PO35 version A2 New PCR mix): hybridization and ligation/amplification of specific probes of each of the 79 exons of the DMD gene and the Dp427c. Capillary electrophoresis on a 3130XL automated sequencer (ABI Prism), analysis of peak heights and peak areas by Gene Mapper software (ABI Prism).  
**Confirmation method:** Semi-quantitative fluorescent multiplex PCR: PCR with fluorescent primer pairs and electrophoresis on a 3130XL automated sequencer (ABI Prism), analysis of peak heights and peak areas by Gene Mapper software (ABI Prism). Target exons: 2, 4. Reference exons: 1, 5, 11, 22  

Please note that DNA has been stored from this patient's sample at this centre, and will be kept indefinitely unless a written request for its disposal is received from the patient or their parent/guardian.  
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**Filename:** D 03988  
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