

# Part 1 examination

## **Genetics: First Paper**

## **Tuesday 25 September 2018**

Candidates must answer FOUR questions

## Time allowed: 3 hours

- 1. How might you evaluate the cost effectiveness of whole exome or genome analysis in clinical practice? Why might you want to do undertake such an evaluation and what would be the challenges?
- 2. Describe the structure, organisation and function of the human genome and use examples to illustrate how this has influenced analysis for the purpose of genetic diagnosis.
- 3. Describe with the use of examples the advantages and disadvantages of different test strategies available for dosage/quantitative analysis (of the genome) for the investigation of clinical genetic disorders.
- 4. You have been asked to write best practice guidance for genetic analysis of ONE of the areas listed below. Provide an outline of the topics to cover and any specific issues to address.

Cystic fibrosis Chromosome 11p15 imprinting disorders Fragile X syndrome Haemaglobinopathies

5. You have been asked by your local Haematologists to give a presentation on current haemato-oncology genetic testing and what will impact on service provision over the next 5 years. Include current and future testing strategies, including the techniques, their advantages and disadvantages.

OR

You have been asked by your local neonatologists to give a presentation on current genetic testing and what will impact on service provision over the next 5 years. Include current and future testing options, including the techniques, their advantages and disadvantages.



## Part 1 Written Examination

### Genetics

## **First Paper**

## Tuesday 26<sup>th</sup> September 2017

#### Candidates must answer FOUR questions

## Time allowed: 3 hours

- 1. Compare and contrast <u>with examples</u> the following FOUR approaches for genetic diagnosis of inherited rare genetic disorders:
  - (i) whole genome sequencing
  - (ii) whole exome sequencing
  - (iii) clinical exome sequencing
  - (iv) next generation sequencing of a targeted panel of genes.

Include the advantages and disadvantages of each approach.

What is the expected impact on service provision for genetic diagnosis of inherited rare genetic disorders over the next 5 years?

- 2. Describe what is meant by epigenetic inheritance. Explain its relevance, <u>with examples</u>, to our understanding of human genetic disorders. Include analytical approaches used for studying such genetic disorders.
- 3. Analysis of tumours for mismatch repair deficiency detects cases with absent mismatch repair protein expression and/or microsatellite instability but no evidence of a germline mutation. Describe the possible explanations for this, how these may be investigated and the clinical relevance.
- 4. Describe <u>with examples</u> the ways in which the utility of a genetic diagnostic test may be assessed. What processes may be implemented to evaluate test performance and use.
- 5. Analysis of cell-free DNA from maternal blood is widely used to provide genetic information in pregnancies. It is proposed that NIPT should be implemented as an additional test within the current Down syndrome screening pathway for high risk mothers. Write a patient information leaflet for NIPT for this purpose. In addition explain why NIPT is a screening and not a diagnostic test.