GeneWatch UK: The Data Protection Bill: Implications for DNA and genetic data

November 2017

GeneWatch UK is a not-for-profit organisation, which aims to ensure that genetic technologies are used in the public interest. This briefing therefore highlights the implications of the Data Protection Bill for the collection, storage and use of DNA and genetic information.

GeneWatch UK welcomes the Government’s decision to introduce the Bill and to seek to implement the provisions of the EU’s new General Data Protection Regulation (GDPR) and Law Enforcement Directive (LED). However, we have a number of concerns about the provisions in the Bill. These relate to the unacceptable inclusion of delegated powers and a large number of overly broad exemptions.

1. Background

Genetic information can act as a “biometric”, may also provide health information about the individual and family members, and in addition reveals a person’s family relationships (Box A).

Due to its sensitivity, genetic information is classed as one of a number of “special categories” of personal data for the purposes of the GDPR and LED. It is widely recognised that it is impossible to anonymise significant amounts of genetic information.

Box A: What is genetic information?

A person’s DNA is contained in every cell in their body and may be extracted from a biological sample (e.g. their blood or saliva) and analysed to obtain genetic information. This can include information about a single genetic mutation (which may occur in many individuals), or multiple genetic differences (which may give a high probability of uniquely identifying an individual), up to an including the whole genome (a long string of chemical letters that is likely unique to a given individual). A forensic DNA profile is a string of numbers based on parts of a person’s DNA which has a high discriminatory power. Genetic information can be stored digitally on a computer database and searched and used like a “genetic fingerprint” to identify an individual. However, unlike a fingerprint, genetic information can also identify a person’s relatives, including non-paternity. Some (but not all) genetic information may provide reveal information about a person’s current or future health and other physical characteristics (such as appearance). This may include information that they themselves may be unaware of (such as the risk of developing a late-onset disease, or of passing a genetic disorder to their children).

Genetic data is currently routinely collected and used by the police, security services, immigration services, commercial companies (for paternity, ancestry and health testing, and for research), universities and research institutions, and the NHS. Currently, genetic testing within the NHS is mainly limited to specific tests in limited patient populations, but there are proposals to expand this significantly in the future, including through public-private partnerships. Some commercial companies already offer “direct to consumer” genetic testing (e.g. the Google-funded company 23andMe, and Ancestry.com). All the major digital companies, including Google, have expressed an interest in storing and analysing genetic data, and Google has repeatedly discussed potential access to genetic information from the NHS with the UK Government.¹
Genetic information has many uses, but is also open to abuse (Box B). To prevent these abuses, and maintain public trust in legitimate uses of DNA and genetic information, it is vital that adequate safeguards are in place.

**Box B: Genetic information: uses and abuses**

Genetic information has important uses in identifying suspects in criminal investigations, resolving paternity disputes, diagnosing genetic disorders, and (in some circumstances) identifying some health risks. However, it is also open to abuse, as stored genetic information can be misused for a range of purposes, including: government surveillance (for example, identifying who attended a political meeting and tracking down their relatives); exposing individual identities and those of their families in ways which may put a person at risk of abuse (for example, by exposing non-paternity in families, or the location of a vulnerable individual); other criminal activity (e.g. identity theft); and commercial tracking and exploitation (for example, targeting persons claimed to be at high genetic risk of disease with marketing for drugs or supplements); genetic discrimination based on interpretations regarding a person's ancestry or further health.

Previous examples of Government failure to maintain trust through adequate safeguards include significant controversy around the retention of innocent people’s DNA profiles on the police National DNA Database (ultimately resolved by the Protection of Freedoms Act 2012)\(^2\); and the furore surrounding Clause 152 of the Coroners and Justice Bill in 2009 (which was rapidly dropped)\(^3\).

2. **Comments on the Bill**

Many of our concerns have already been highlighted by others, including Liberty\(^4\), Privacy International\(^5\), and MedConfidential\(^6\). The focus of this briefing is to highlight why adequate safeguards are particularly important in the context of genetic information.

2.1 **Delegated powers**

There is significant concern regarding the powers given to the Secretary of State in Clause 15 and Clause 9(6 and 7). These clauses allow the Government wide powers to alter the application of the GDPR and to amend Schedule 1 (conditions for processing special categories of personal data, which includes genetic data).

Similarly, Clause 33(6) provides delegated powers to amend safeguards for processing of sensitive personal data for law enforcement; Clause 84(3) provides delegated powers to amend safeguards for processing of sensitive personal data by intelligence services; and Clause 111 provides delegated powers to make further exemptions from data rights regarding intelligence services processing.

In GeneWatch’s view these clauses make the protections in the Bill meaningless as they remove any requirement for detailed parliamentary scrutiny, should the Government choose to weaken or even totally remove the safeguards in the Bill. In order to maintain public trust that genetic information and other data are adequately protected by law, these clauses need to be deleted.

2.2 **Automated decision-making**

Clause 13 provides for automated decision-making authorised by law in some circumstances. Similarly, law enforcement agencies are exempted from the prohibition on making purely automated, significant decisions by clauses 47 and 48, and intelligence agencies by clauses 94 and 95. However, these clauses do not provide for any safeguards.
to prevent discrimination, or to ensure that the decisions are based on a valid analysis of the available data.

Genetic information may inform decisions regarding: the identity of an individual or the relatedness of families (e.g. for immigration or maintenance payment purposes); the pursuit of an investigation of an individual, or their family member (using ‘familial searching’ for partial DNA profile matches), for a criminal offence; or the diagnosis or prediction of disease. It is likely that such decisions may become more automated in future, through the use of computer algorithms. In the health area, in particular commercial black-box algorithms may provide highly misleading information about a person’s health. There are also many examples of misidentification of criminal suspects using genetic databases. In some cases, commercial computer software has been used to try to identify an individual’s DNA profile from a mixture: however such results are always open to interpretation. It is important that any algorithms can be independently validated and that their limitations are known to the police and to the courts.

Further safeguards are therefore essential to: (i) ensure the analysis of genetic data is valid and undertaken in an appropriate context (with expert involvement as necessary); and (ii) prevent genetic discrimination.

Some relevant safeguards already exist: for example, Crown Prosecution Service guidelines require corroborating evidence if a DNA match is used in a prosecution; and the Equality Act 2010 restricts what employers can ask about in pre-employment medical checks, which limits the potential for genetic discrimination. However, the status of other protections post-Brexit remains unclear. For example, the Charter of Fundamental Rights of the European Union prohibits discrimination based on genetic features (Article 21), but no such prohibition exists in domestic law. The Forensic Regulator lacks statutory powers to enforce standards, including for DNA collection and analysis. The In Vitro Diagnostics Regulation (IVDR) requires clinical evidence to support claims regarding the diagnostic and predictive value of medical tests, including genetic tests: but the IVDR is not operational for the purposes of the EU (Withdrawal) Bill, and therefore will not be automatically incorporated into UK law.

GeneWatch UK believes it is premature to add broad exemptions to the Bill to allow for automated decision-making, as safeguards to ensure valid data analysis and prevent discrimination are currently inadequate.

**2.3 National Security, Defence and Immigration**

Numerous broad exemptions to the rights of individuals in relation to processing for national security, defence and immigration purposes have been highlighted by Liberty, Privacy International and the Information Commissioner’s Office.

Rapid DNA analysis, which allows on the spot automated DNA testing for policing purposes, is beginning to be introduced. This speeds up the process by avoiding the need for a sample to be sent to experts in a laboratory, but it also raises concerns that on-the-spot testing might in future be used in the streets or at borders, posing major risks to human rights. Such uses could fall within the exemptions as proposed, posing an unacceptable risk to the rights of individuals.

Data protection or privacy laws which cover police information, including genetic information, are an important part of international best practice worldwide, and governments increasingly require standards to be met before sharing genetic information across borders for policing purposes. Similar safeguards are increasingly likely to be required for cooperation in intelligence and border control: not least because the expected increasing volumes of data
will increase the number of mistakes. Thus, failure to provide adequate safeguards could not only damage public trust but also prevent legitimate cross-border co-operation.

GeneWatch UK therefore believes that any exemptions for national security, defence and immigration purposes should be strictly limited to prevent abuses.

References

8. DNA database in doubt after teenager spends three months behind bars for rape in city he has never even visited because gene samples were mixed up. Daily Mail. 18th May 2012. http://www.dailymail.co.uk/news/article-2114252/Teenager-spends-months-bars-DNA-blunder-fingers-rape-city-visited.html