DRAFT REPORT OF THE IBC ON BIG DATA AND HEALTH

Within the framework of its work programme for 2016-2017, the International Bioethics Committee of UNESCO (IBC) decided to address the topic of Big Data and health, including but not limited to the issues of autonomy, consent, data protection, governance, etc.

At the 22nd (Ordinary) Session of the IBC in September 2015, the Committee established a Working Group to develop an initial reflection on this topic. The IBC Working Group, using email exchanges, started preparing a text on this reflection between October 2015 and May 2016. It also met in Cologne in May 2016 to refine the structure and content of its text. Based on this work, the IBC Working Group prepared a preliminary draft report which was discussed during its 23rd (Ordinary) Session in September 2016. As a follow-up to this discussion, the IBC Working Group started to revise the preliminary draft report between September and December 2016. The IBC Working Group met in Spain in March 2017 to further refine the text. This document contains the revised text in the form of a draft report, and will be submitted to the IGBC, the IBC, and COMEST between May and June 2017 for comments.

As it stands, this draft report does not necessarily represent the final opinion of the IBC and it is subject to further discussion within the Committee in September 2017. This document also does not pretend to be exhaustive and does not necessarily represent the views of the Member States of UNESCO.
DRAFT REPORT OF THE IBC ON BIG DATA AND HEALTH

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I. SCOPE AND DEFINITIONS

1. The digital form of all kinds of data is leading to an exponentially evolving phenomenon called Big Data. It is touching and transforming every area of human life all over the world. In this report the IBC focuses on issues relevant to the area of health on the individual and the public level. It presents recommendations so that the full potential of Big Data can be tapped while at the same time human dignity, fundamental rights and fundamental freedoms are upheld according to Article 3 of the Universal Declaration on Bioethics and Human Rights (2005).

2. Big Data is characterized by the so-called 5 Vs:
   a. **Volume** refers to the huge amount of digital data. It is growing exponentially. While three-fourths of data were analogous in 2000, today more than 99% of all data are digital data. For 2020 there is an estimated amount of 44 zetabyte \(10^{21}\) digital data, for 2025 it is 180 zetabyte (IDC, 2014). But volume of data alone, e.g. coming from whole genome sequencing, does not already constitute Big Data.
   b. **Variety** hints at the fact that there are different kinds of data from diverse kinds of sources. For healthcare and research several sources are relevant: medical data from individual patient care, public health data, data from different insurances, research data which are collected by researchers and citizen scientists, companies or individuals themselves, life-style data e.g. from health apps, data from social networks and data from commerce. These data can be classified in different ways and according to different criteria: there are e.g. personal data, anonymized data, metadata, primary and secondary data.
   c. **Velocity** means the very high speed at which data can be collected and processed. Real-time-tracking and cloud solutions allow for comprehensive processing within seconds, even producing recommendations e.g. for medication, behavior or nutrition.
   d. **Validity** refers to quality of the data and the question if they really show what they are meant to show regarding content and precision. The context of data plays a major role here.
   e. **Value** finally draws attention to the meaning of the data for a specific question e.g. with regard to a certain disease. Here again it is important to take the context of data into account.

3. Against this background the IBC uses the term Big Data in the area of health as referring to large collections of complex health-related data sets from multiple sources that cannot be processed with traditional applications, but require development of new algorithms and enhanced computing power, often in real time. Typically such data sets cover very large numbers of individuals, but analysis of all available data from one single patient under certain conditions can also be considered Big Data analysis.

4. There is no all-agreed definition of health which could be applied in every context of application to every period in life course and in every area of the world. Definitions rather vary according to perspective (a subjective understanding or objective concept) and according to the purpose of the definition in terms of therapeutic or preventive actions which then can be legitimated. For example, the broad definition of the World Health Organization (WHO) is meant as a regulative leading idea to foster global health. It defines health as the state of complete physical, mental and social well-being and not merely the absence of

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1 In this report the term citizen science refers to research projects which are conducted by lay people, often cooperating or being lead by professional scientists or research institutions.
disease or infirmity, thus embracing the whole life of an individual in every respect. Compared with this a narrower approach is preferred in national health care systems, specifying and limiting the responsibilities of medical professionals and institutions. Especially, the WHO definition shows that everything in life matters with regard to health, thus suggesting a Big Data approach to health care and research.

5. In times of information and communication technologies (ICTs) new terms like eHealth (electronic health) and mHealth (mobile health) have come up. WHO defines eHealth as “the use of information and communication technologies for health” (WHO, n.d.), and mHealth as a subcomponent of eHealth in terms of “medical and public health practice supported by mobile devices, such as mobile phones, patient monitoring devices, personal digital assistants (PDAs), and other wireless devices” (WHO, 2011).

6. There is a day-to-day increasing number of apps (according to estimations there are already more than 200,000) which address health related issues. Regardless of the impossibility to control and to guarantee the quality of all these apps it is hardly possible to sharply delineate medical from nonmedical health apps. They mainly fall into three categories: firstly treatment-related apps, secondly prediction and prevention related apps, and thirdly life-style related apps. Accordingly, when addressing Big Data and health in this report, this necessarily entails more than traditional health care and health research.

II. LEGAL REGULATION

7. There are no specific regulations of the phenomenon of Big Data in the national and international legal frameworks. Nevertheless, there is a complete regulatory framework for personal data protection in many legal jurisdictions, mainly in Europe, of which many rules can be applicable in the area of Big Data, though it is a new reality in the sense of quantity, analysis, accessibility, and application. Furthermore, countries that have no specific laws on data protection can still use constitutional and statutory law provisions as well as common law principles for the same purpose. A good example in this regard is most commonwealth countries (UNCTAD, 2016). So there is not a lack of regulation but of specific provisions and perhaps of new principles which are adequate to regulate the new features of Big Data.

8. In the international legal framework, Article 12 of the Universal Declaration of Human Rights, adopted by the United Nations (UN) General Assembly in 1948, covers privacy, stating that: “No one shall be subjected to arbitrary interference with his privacy, family, home or correspondence, nor to attacks upon his honour and reputation. Everyone has the right to the protection of the law against such interference or attacks” (UN, 1948). In a similar sense, Article 8(1) of the European Convention on Human Rights states that: “[e]veryone has the right to respect for his private and family life, his home and his correspondence” (CoE, 1950). Article 8(2) adds that: “[t]here shall be no interference by a public authority with the exercise of this right except such as is in accordance with the law and is necessary in a democratic society in the interests of national security, public safety or the economic well-being of the country, for the prevention of disorder or crime, for the protection of health or morals, or for the protection of the rights and freedoms of others” (CoE, 1950).

9. Many of the regulations at the international level have been developed in the context of international data flows mainly due to trade in health services, which leads to cross-border data transfers. The United Nations Conference on Trade and Development (UNCTAD)’s Data protection regulations and international data flows (2016) is notable in this regard. Agreement on the core principles can be attributed to the United Nations General Assembly’s Guidelines for the Regulation of Computerized Personal Data Files (1990), which contain principles to ensure protection of privacy and confidentiality that as a minimum must be provided for in national legislations. These are the principles of purpose-specification and security. The guidelines equally require countries to designate an authority that supervises the observance of these principles, sanction those in breach and prescribe the need to protect privacy during the trans-border movement of personal data. The
guidelines were meant to govern computerised and manual files that contain personal information (see para 10 of the guidelines) but the principles can still be applied, to some extent, in the context of Big Data. Other non-legally binding guidelines, which have shaped national legal frameworks, are the World Medical Association’s Declaration of Taipei on Ethical Considerations Regarding Health Databases and Biobanks (2016) and the Declaration of Helsinki (2013). While there is a broad consensus on the core data protection principles at the heart of most national laws and international regimes, the main challenge is divergence in the implementation of these principles as well as in the detailed data protection laws of the world (UNCTAD, 2016).

10. The Organisaton for Economic Cooperation and Development (OECD) has developed its Guidelines Governing the Protection of Privacy and Transborder Flow of Personal Data (2013). There is also an OECD report which highlights the need for development of appropriate legal frameworks for sharing information (OECD, 2010). This report stresses the need for new legal framework which allows for sharing of health related information between health care professions within and across health care organizations, as well as across organizational and geographical boundaries. The report notes that very few countries in its remit have really addressed these challenges. More recently, in January 2017, the OECD has published its Recommendation on Health Data Governance. They highlight the importance of complementing legal data protection through education and awareness raising, skills development, and the promotion of technical measures. The Recommendation calls upon countries to develop and implement health data governance frameworks that secure privacy while enabling health data uses that are in the public interest in accordance to twelve high-level principles.

11. With regard to all regulations, an important distinction has to be made between Europe and the US. The European approach is based on a view that privacy is a fundamental human right and it involves top-down regulation and the imposition of across-the-board rules restricting the use of data, or requiring explicit consent for that use. The United States, in contrast, employs a sectoral approach that focuses on regulating specific risks of privacy harm in particular contexts, such as health care and credit. This places fewer broad rules on the use of data, allowing industry to be more innovative in its products and services, while also sometimes leaving unregulated potential uses of information that fall between sectors. (USA, 2014)

12. Different approaches in the two regions led to the design of the EU-US and Swiss-US Privacy Shield Frameworks (the privacy shield) by the US Department of Commerce, the European Commission and Swiss Administration for purposes of ensuring compliance with personal data protection requirements in transatlantic commerce (USA, n.d.). The privacy shield replaced the previous Safe Harbor Agreement of 2000 and is subject to annual review to ensure its currency as technology changes and the EU data protection regime is transformed (cf para 27 below) (Weiss and Archick, 2016).

13. The Council of Europe approved in 1981 the Convention for the Protection of Individuals with regard to Automatic Processing of Personal Data. The aim was to protect the individual against abuses which may be associated with the collection and processing of personal data. Later, the Council of Europe approved an Additional Protocol to this Convention (CoE, 2001) regarding supervisory authorities and transborder data flows. It provides for the setting up of national supervisory authorities responsible for ensuring compliance with laws or regulations adopted in pursuance of the Convention. Furthermore data may only be transferred if the recipient State or international organization is able to provide an adequate level of protection.

14. The EU has only a limited legal competency on health matters, which can be used mainly to promote cooperation and coordination among Member States. However, in the area of data protection there is a common regulation through Directive 95/46/EC of the European Parliament and of the Council of 24 October 1995 on the protection of individuals
with regard to the processing of personal data and on the free movement of such data (EU, 1995). This situation will change in 2018. Instead of a Directive there will be a Regulation which is directly binding in all Member States and is relevant for extraeuropean companies as well (EU, 2016).

a. The new Regulation provides that the processing of personal data should be designed to serve mankind. The right to the protection of personal data must be considered in relation to its function in society and be balanced against other fundamental rights. This Regulation respects all fundamental rights, freedoms and principles recognized in the Charter as enshrined in the Treaties, in particular the respect for private and family life, home and communications, the protection of personal data, freedom of thought, conscience and religion, freedom of expression and information, freedom to conduct a business, the right to an effective remedy and to a fair trial, as well as cultural, religious and linguistic diversity.

b. In relation to consent, the Regulation establishes that it should be given by a clear affirmative act. If the data subject's consent is to be given following a request by electronic means, the request must be clear, concise and not unnecessarily disruptive to the use of the service for which it is provided. It later adds that “[c]onsent should not be regarded as freely given if the data subject has no genuine or free choice or is unable to refuse or withdraw consent without detriment” (EU, 2016; para 42).

c. In the area of research, the Regulation comments that, provided that recognised ethical standards for scientific research are kept, data subjects should be allowed to give their consent to certain areas of scientific research, since the specific purpose of personal data processing can often not be fully identified at the time of data collection.

d. In the area of public health, the new Regulation mentions that the processing of special categories of personal data may be necessary for reasons of public interest without consent of the data subject. Such processing should be subject to suitable and specific measures to protect the rights and freedoms of natural persons.

15. The United States of America offer also a legal framework through some recent Acts. There are no regulations that concern Big Data but companies undertaking Big Data processing operations in the area of health need to comply with data protection regulation at the federal level: the Health Insurance Portability and Accountability Act (HIPAA) and the HIPAA Privacy Rule (USA, 1996; USA, 2002). Both regulations require appropriate safeguards to protect the privacy of personal health information, and set limits and conditions on the uses and disclosures that may be made of such information without patient authorization. The rule also gives patients rights over their health information, including rights to examine and obtain a copy of their health records, and to request corrections. The American Recovery and Reinvestment Act (ARRA) and the Health Information Technology for Economic and Clinical Health Act (HITECH) create significant incentives for an expanded use of electronic health record and they also contain some rules about data protection, mainly about measures in relation to breaches of health information (USA, 2009).

16. The African Union (AU) and the Asia Pacific Economic Cooperation (APEC) are other examples of regions that have developed regional legal frameworks for the protection of personal data and privacy. The AU’s Convention on Cybersecurity and Personal Data Protection requires the processing of personal data involving genetic information and health research to be undertaken with the authorisation of the national protection authority (AU, 2014; Article 10[4][a]). The updated APEC Privacy Framework provides for “a multilateral mechanism which enables Privacy Enforcement Authorities in the APEC region to cooperate in cross-border privacy enforcement of Privacy Laws” (APEC, 2015).
17. With regard to the protection of data, autonomy and privacy, the legal concept of ownership is important as well. It has to be clearly distinguished between what constitutes a discovery and what constitutes an invention. Normally, new inventions are patentable, as they entail an inventive activity and are used industrially. On July 3, 2012, the European Court of Justice (ECJ) published its landmark decision in Used Soft GmbH v Oracle International Corp (C-128/11). The decision implies that there is specific ownership attributed to intangible goods like software downloaded via the Internet. Although the applicability of this model to other digital goods remains to be considered in future court decisions, the ECJ has opened the door for a discussion on ownership of intangible assets (Hoeren, 2014). Furthermore in an appeal of the judgment in Football Dataco Ltd v Brittens Pools Ltd, April 2010, the European Court of Justice (ECJ) emphasized that the purpose of the Database Directive is to: “stimulate the creation of data storage and processing systems in order to contribute to the development of an information market … and not to protect the creation of materials capable of being collected in a database”.

18. At the international level the TRIPS agreement, Article 10 (2), protects compilations of data or other material in either machine readable or another form. For such data or material to qualify for copyright protection, the Article prescribes that “the selection or arrangement of their contents [must] constitute intellectual creations …” (WTO, 1994). The Article further stipulates the scope of rights in such compilations by explicitly providing that “such protection, which shall not extend to the data or material itself, shall be without prejudice to any copyright subsisting in the data or material itself” (WTO, 1994).

19. The Council of the European Union has developed important statements about data base rights (EU, 1995; EU, 1996 (Article 7, paragraph 4)). For example: specific and separate legal rights last for 15 years, but each time a database is substantially modified, a new set of rights is created. An owner has the right to object to the copying of substantial parts of his database, even if data is extracted and reconstructed piecemeal. The arrangement, selection, and presentation of the data may be protected by copyright, while the database right can protect a whole database.

III. VISIONS, TRENDS AND CHALLENGES

20. One vision of Big Data in health care is a comprehensive and evidence-based personalized or so-called precision medicine, which combines the best available scientific knowledge with personal experience of health professionals for the benefit of the individual patient. It is based on technological developments in genomics and other high-throughput “omics” techniques which have made molecular analysis of human samples orders of magnitude cheaper and more efficient. This molecular data can now be complemented with digital imaging data spanning from the microscopic level to whole body imaging and with environmental and lifestyle information collected from a large number of individuals (e.g. population or patient cohorts), from surveys or from different registries, databases and research infrastructures. Furthermore data can be collected on social environment, communication and behavior, thereby bringing us closer to a more comprehensive understanding of health and disease according to the biopsychosocial concept as put forward by the WHO. According to this vision, all this data combined with information in the electronic health records (EHR) will in the future provide a fundamentally different approach to diagnose and treat patients in a personalized way, i.e. to offer the right recommendations and individually tailored treatment for a person at the right time. It also can foster patient safety, combining different data from different sources in order to analyze and eventually prevent adverse events.

21. A related vision is that having all this in-depth knowledge of an individual will gradually make it possible to determine his/her predisposition and risk profile to develop a disease and to deliver timely and targeted advice for prevention.
Furthermore it is likely that in the future health care will move more and more towards remote collecting data for diagnosing, monitoring and supporting therapy. Collected data can then be used to detect early warnings of disease like an imminent heart attack and make recommendations for adequate behavior. It can also contribute to improved telemedical health care for people living in remote regions. This can allow better access to quality health care and thereby contribute to global health.

Patients can have the possibility of greater access to and control over their data, e.g. having their EHRs in their smartphone. The smartphone could actually turn out to be a central device for coordinating one’s own health care and for creating one’s own health network thus democratizing medicine (Topol, 2015), while fostering autonomy and health literacy.

Having better access to health information as well as individualized profiling and recommendations – provided they are quality approved – citizens will potentially profit from better understanding their health status and health improving behavior. However, there is a human-dependent challenge: the fact that even if one understood the information it does not necessarily mean acting accordingly, and in fact, often no health-promoting action whatsoever is taken. Tobacco smoking is a striking example.

Against this background at least four paradigm shifts in individual health care are likely to occur: a shift from disease orientation to health orientation, from focus on therapy to prevention, from health to life-style counseling and from the role of a patient to the role of a user, customer or digital citizen.

For the pharmaceutical and medical device industry there is the hope that Big Data will foster the understanding of diseases and their underlying mechanism and thereby lead to development of new targeted drugs, devices and treatments. Such Big Data is also expected to help design stratified clinical studies thereby reducing the number of participants and costs, and give quicker results.

But these visions of Big Data do not only pertain to individual health care and research. Big Data approaches are also supposed to provide a lot of new information in order to strengthen the evidence base for public health policies, e.g. enabling better risk-adjusted prevention strategies for defined target groups.

Regulators might better understand and control study designs and their policies might benefit from improved pharmacovigilance. Once a new drug has entered the market, Big Data allows for real world data collection and assessment in a large number of patients over a long period of time. As the majority of the global population lives in areas covered by mobile cellular networks, the number of citizens who can contribute data is bound to increase.

Furthermore Big Data can contribute to support learning health care systems (IOM, 2007; IOM, 2013). By analyzing real world data from health care in a structured and quality controlled way everyday experiences can inform the best and most efficient way to administer diagnostic, therapeutic and preventive measures as well as shaping structures for health care and research. Examples are the work on learning health care systems of the OECD, EU projects such as the Coordinated Research Infrastructures Building Enduring Life-science Services (CORBEL) Project (http://www.corbel-project.eu/about-corbel.html), and practical instruments that facilitate data sharing such as Innovative Medicines Initiative (IMI), and industry initiatives (OECD, 2013b; OECD, 2015; IMI, 2014; PhRMA, 2014). Recently, the European Medicines Agency (EMA) published its guidance on the release of anonymised individual patient (trial) data (EMA, 2016).

Having in mind the broad definition of health by the WHO and recognizing that the major part of an individual’s health status does not depend on health care but on social determinants, including but not limited to education, life style, and environment, Big Data opens up the way to a holistic view on health by bringing together different kinds of data e.g.
from registries, apps and health records. As a consequence, the line between health and life style issues in health care becomes increasingly blurred.

31. These visions and trends are accompanied by major challenges which are of a technological as well as of an ethical, social and legal nature.\(^2\)

32. Welcoming the possibility of a holistic view on health the line between the health care sector and other societal sectors is increasingly blurring at the same time. In addition to the traditional sources of health data, such as medical records and laboratory results, other sources which are not traditionally regarded as health related are used, for example social networks or consumer data, public data sources from non-health areas such as the Internal Revenue Service, Education or Social Services Departments. Search engine providers also request and collect much information on their users, which they later process and sell to different companies which in turn use “personalized marketing strategies”, offering users different promotions based on their search histories or participation in online groups, for instance. Such customer profiling also includes health issues – a simple “private” internet search on a personal or familial condition becomes public information. So obviously, there are complex challenges to data protection and privacy as well as to the quality of data.

33. Machine-learning prediction methods have been very productive in medicine. However it is essential to understand underlying assumptions and ensure that conditions like stability (namely the conditions in which the data were collected remain the same) are met to secure the quality of data as well as the validity and usefulness of the conclusions. Big Data mainly allows for finding patterns and correlations. These can be misunderstood as causal relationships so that inappropriate and harmful consequences can emerge. So, appropriate skills and competences by all involved actors are a major challenge as well. Health professionals will have to understand and to properly verify and apply available data. Researchers, citizens and policy makers have to understand the relevance of different contexts of data and the implications of algorithms. This requires transparency and interpretability of data and their contexts. This only is possible if algorithms are accessible, and if experimental design and optimal resource allocation policies are given.

34. The line between different professional disciplines will also blur. At least some diseases will no longer be understood according to the affected organ, but rather according to the underlying molecular mechanism, the pattern of mutations and variants in the “omics”. A multidisciplinary approach for health care and research, as is already implemented in oncology in many countries, is needed to optimize the treatment of individual patients which may also result in new professional disciplines. Furthermore the integration of disciplines like social science, behavioral economics, epidemiology etc. from the beginning seems to be adequate in view of a comprehensive understanding of health as well as of the use of data in causal inference methods.

35. Eventually Big Data and an environment full of sensors lead to the effect that individuals become more and more transparent while the technological environment, algorithms, as well as the consequences of data analysis and the underlying weighting of factors become increasingly opaque. This leads to several ethical challenges especially with regard to autonomy, privacy and justice (see Chapter IV).

36. In many countries, particularly in the developing world the ethical and scientific infrastructure for research are still developing and many are still far from harnessing the potential benefits that the use of big data promises. Access to data remains a challenge and the health information platforms that can support the storage of huge amounts of data and to translate existing data into real action are still lacking. This calls for some attention to capacity building. Capacity strengthening is an important way of ensuring that developing world scientists and health professionals do not become just data collectors in international

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\(^2\) For further detail, especially on ethical aspects, see Chapter IV.
collaborative projects but active participants in transforming data into tangible health benefits as a form of reciprocity.

37. Furthermore such advanced health care towards precision medicine is expensive and limited to richer countries and Big Data approaches do not yet cover diseases affecting people in countries with lower health-care spending. To implement Big Data-driven precision medicine new solutions for data handling in actual health care settings are needed. Only then could Big Data be used to improve medical interventions and health services, prediction and prevention strategies and health policies in general.

38. It is very difficult to foresee how fast and to what degree the trends described above will be made a widespread reality. Despite the sketched optimistic vision, it is important to be cautious and to avoid exaggeration of the current state of scientific knowledge and the potential benefits of Big Data and precision medicine for health care globally. The "hype of Big Data" can lead to overstatements and unrealistic estimations, to unbalanced health care policy priorities and marketing of unproven products and services. On the other hand there is the danger of losing the benefits of Big Data by neglecting its potentials. A balanced way of dealing with hopes and promises is very important.

IV. ETHICAL ASPECTS

IV.1. Autonomy and Consent

39. Autonomy of an individual in terms of exercise of his or her self-determination includes seven dimensions:

a. Competence of the individual concerned to access, to understand, to assess and to apply relevant information;

b. Information has to be available that is relevant and understandable for the question at stake;

c. There has to be a choice between different options, be it the one between doing or abstaining from doing something or the one between doing different things;

d. Values of the individual, his or her preferences and attitudes are taken into account in deciding and acting;

e. Voluntariness is granted so that the individual can decide and act without inner or outer coercion;

f. Formation of will refers to the ability of the individual to choose an aim and appropriate means to reach it;

g. Action can mean a conscious doing or refraining.

40. The main guarantee, which has traditionally been established to protect the autonomy of human subjects in health care and research, is consent. Autonomy and responsibility as well as consent and protection of persons without the capacity to consent are addressed in Articles 5, 6 and 7 of the Universal Declaration on Bioethics and Human Rights (UDBHR). Article 6 provides that any preventive, diagnostic and therapeutic medical intervention as well as scientific research should only be carried out "with the prior, free, express and informed consent of the person concerned" (UNESCO, 2005). The Nuremberg Code concentrates on consent and is at the origin of the concept that participation in research is a voluntary activity. The Declaration of Helsinki also enshrines consent as a main guarantee.

41. The scale and complexity of personal data processing in the area of Big Data make it virtually impossible for people to keep track or make meaningful decisions about all uses of their personal data. There are major problems with regard to all seven dimensions which are mentioned above. Even if individuals could keep track and would be informed, their
decisions might be skewed by various decision-making difficulties. This leads to a loss of autonomy which in consequence means a loss of control and a loss of freedom from decisions which are made by the technological environment and automated processes. Governance has to compensate for this loss of control, ensuring that all the dimensions of autonomy are respected and supported (see Chapter V).

42. A further problem of informed/informatic consent is the impossibility to identify the age of the user, and so to tailor the information to be presented according to his or her age and capability of understanding (EC, 2012). The education of minors as active users of these new technologies is particularly important.

43. In the context of clinical trials, consent has traditionally been specific, that is, subjects are asked to consent to a specific research project. In many such cases, the consent obtained from research participants does not extend to the use of samples or data outside the original or primary research they have consented to. Valid consent requires that adequate information is disclosed and that potential subjects understand the nature, risks and potential benefits of the research and thus grant voluntary informed consent.

44. In the area large databases like biobanks, and even more in the area of Big Data research, the potential use of data in the future cannot be predicted. It also involves interrelationships between multiple and changing data sources, both medical and non-medical. This has led to calls for a new, more appropriate model of consent that allows a wide use of data while at the same time respecting the participant’s or patient’s autonomy. Already the field of biobanks has put forward new suggestions for consent provision such as what is called broad consent.

45. Broad consent can cover a wide range of activities while at the same time being narrowed down to limited, though just vaguely specified uses such as research into the causes of complex diseases. Thus broad consent is not the opposite of “specific” consent because according to the understanding of the IBC it is no blanket or open consent, which does not have any conditions attached. It just typically operates at a higher level of abstraction. Broad consent leaves the essential informed consent model intact, but an individual simply consents to a range of possible research that could be done with her/his information in relation to a specific area or line of investigation. The conditions for broad consent to be valid is the need for some form of governance where e.g. a relevant committee reviews a proposal to ensure that the rights and interests of individuals are adequately protected.

46. This formula of informed consent is getting more and more accepted (WMA, CIOMS), and is utilized in Europe, where individuals broadly consent to all possible research with their tissue related to the same or similar area or line of research. It is also a model that is gaining attention in developing countries, particularly in the context of genomics and biobanking in Africa.

47. Another widely accepted approach is opt-out consent according to which health data can be used for research purposes unless an individual affirmatively opts out. This approach requires adequate information beforehand, address the potential implications of permitting biospecimens and personal data to be collected. So it requires practitioners to provide adequate education and information although time often is scarce. Differing from the research setting the virtual impossibility to opt-out from the commercial health-related databases (of mobile phone and internet operators, apps, commercial and consumer behavior databases) is a particular concern, as there is virtually no control over such databases.

48. According to the principles of participation and transparency a further model which is mainly applicable because of widespread ICTs has been proposed. It is called “dynamic consent”. Based on an initial consent from the participant it entails an update on the use of data in a continuous way so that the individual can opt out for specific uses of data while
allowing the use for other purposes. It requires the involvement of public powers both as a
guarantee of the individual rights as well as the promotion of individuals’ participation
through education and information. It calls for empowering subjects and patients to be able
to monitor the use of their health data. Dynamic consent could in some settings allow for
control of data access by individuals, enabled by mechanisms such as consent portals. As
the Nuffield Council in the UK has pointed out, continuing participation can have the
advantage of allowing participants to shape the possibilities of research through their
decisions about what uses of data to permit by effectively “voting” for those uses by
consenting to them. The individual effectively participates in the enterprise. In a way, the
project becomes a shared or joint enterprise of the patient along with the researcher, and so
is not being used or exploited. While this model of consent could work well in developed
countries with advanced technologies that allow patients to monitor how their data is being
used through a database system, and with time and willingness to devote it to follow the use
made of their data, its implementation in less developed settings could face several
challenges such as the lack of local technologies and low literacy rates that could hinder
patients’ comprehension in the context of health care and participation in research.

49. Consent is a major challenge with regard to health apps and in the area of social
media and networks. The digital information is generally written in small characters to be
displayed on smartphones, sometimes with no real alternative of dissent. The process of
giving informed consent on screen (and not on paper), in a setting where the digital user
does not have access to consultation, often leads to clicking in an immediate and fast way
without sufficient time for making an informed choice and without the possibility of
ascertaining actual awareness, competence and voluntariness. Trade terms, which are
usually too long and for most people hardly understandable, are seldom read at all. In this
sense it is doubtful that the “one click, accept all”-procedure is really a free consent.

50. Facing the challenges for obtaining an autonomous consent for the collection and
use of big data and the limitations of all proposed models of consent, it is important that
autonomy is supported by public education which allows users, patients and research
participants to understand the range and the impact of possible uses of their data.

51. Though it must be stressed that autonomy links directly with a person’s dignity, there
are also major collective interests which support research being carried out in the field of Big
Data and health in order to promote evidence-based precision medicine as well as learning
health care systems. Therefore, a balance between individual and collective interests must
be sought. Thus a solidary claim to consent to participation in databased health research by
contributing one’s anonymized – not personal – data for large databases can be discussed.
According to a principle of solidary reciprocity an individual who profits from former health
research can be supposed to have an interest that other people can profit from progress in
research as well – at least when everyone can be sure that the outcome is beneficial in
terms of a democratically defined public good, and when misuse and unlegitimated
deanonymization of data are strongly prosecuted and punished. In such a framework, there
are at least two main elements which could be useful to balance the individual and collective
interest: firstly the vulnerability of the subjects from whom data are collected; secondly, the
purposes for which the data will be used.

IV.2. Privacy and Confidentiality

52. According to Article 9 of the UDBHR, the privacy of the person concerned and the
confidentiality of their personal information should be respected (UNESCO, 2005). Such
information should not be used or disclosed for purposes other than those for which it was
collected or consented to. In times of Big Data, such protection of privacy and confidentiality
is facing several challenges.

53. Traditional data protection principles such as purpose limitation, data scarcity and
minimization, special protection of sensitive data, fair processing, and safeguarding the
rights of the persons concerned have been widely implemented in order to protect privacy,
though there are different protection schemes and underlying concepts in different regions of
the world. By way of contrast, Big Data inherently entails change and openness of purpose,
limitlessness of data, intransparent processing, little protection of data in order to gain as
much knowledge as possible, and intransparency for persons concerned. This is especially
true with regard to algorithms getting more and more complicated and incomprehensible in
times of artificial intelligence and deep learning.

54. Furthermore anonymization of personal data no longer provides sufficient protection
of privacy and confidentiality. By integrating large amounts of data from different kinds of
sources the reidentification of the person concerned basically becomes possible.

55. In addition more and more differentiated profiling is done, which allows for acting on
groups according to specific profiles (the so called “group identity”). Thus privacy of the
individual is no longer adequately protected even if the individual’s name is not known, but
e.g. the IP address of his or her smartphone or computer. Thus group privacy is an
increasingly pertinent concept in Big Data applications.

56. In times of Big Data, privacy as a concept has shown to be more than mere data
protection. When collecting, storing, processing, analyzing, and applying data which then are
collected, stored etc. again and again (the so called cybernetic loop) it also encompasses
the intrusion into the private sphere by informations, offers, advertisement and the like on
the screen of electronic devices according to one’s personal profile which is constantly
assorted by intransparent algorithms on the basis of a variety of data from diverse sources.
This intrusion touches several dimensions of private life like personal communication,
location, and participation in associations.

57. Eventually the right to privacy is part of the right to freedom with its diverse aspects
like freedom of speech, of association, of location and space, of beliefs, thoughts and
feelings, as well as of behavior and action. Against this background, the IBC uses the term
privacy in the sense of a right to respect for private life in relation to those areas of life or
those data that individuals want to keep reserved for themselves or, at least, for some
specific members of their families or relationships.

58. While individuals have privacy interests, they also share collective interests in the
wider use of data for health research. This broader public interest may come into conflict
with individual privacy. But the relationship between privacy and public interest is not simply
one of opposition. The two are mutually implicated in each other: there are private interests
in the achievement of common goals and a public interest in the protection of privacy that
encourages cooperation. This complex relationship leads to a need to reconcile the
articulation of the private within the public and the public within the private.

59. From several surveys we know that people are aware of having little control over
collection and mining of their data although a majority of them wants to have a certain
degree of control. In the long run this might lead to a lack of trust. The loss of trust might well
cause serious damage to future essential endeavours and projects to foster public health. In
order to protect privacy and the public good at the same time, it is essential that the
population is able to trust in the good use and protection of their health data through a mix of
approaches like law, governance, public surveillance, privacy by design of a device or an
app and privacy by default.

60. New models of participation in data collection and mining, as outlined in the chapter
on autonomy, proper public education on the implications of the use of Big Data, effective
data protection management, accountability of use to both participants and society in
general, as well as innovative models of data ownership and trusteeship are to be developed
in order to protect privacy in the here outlined wide understanding.

IV.3. From ownership to custodianship and benefit sharing
61. Ownership of personal data and the rights entailed, among which we might highlight the restriction of access by third parties, must be recognized, without any doubt, as being of the person/group from whom the data derived. Ownership grounds the empowerment of data subjects to track and check the existence and manipulation of their personal data. Maintaining control of what we are, what we do and what we think and what we present to others and society in terms of personal information is a key facet of liberty in the 21st Century (see also Chapter IV.2).

62. In the area of Big Data, it is important to distinguish between two different concepts linked to the idea of ownership, as it is done in similar areas like biomedicine: that of ownership of personal data and ownership of operational results, ownership as a mechanism to control data and ownership in order to refer to owning products like algorithms, drugs or monitoring tools etc., and intellectual property.

63. It is argued that the benefits of data will be harnessed through fair data sharing practices with the wider scientific community and that this process will facilitate scientific discovery. There is also growing scholarship on the appropriate ethical frameworks that should guide best practices in data sharing in international research collaborations. This includes the use of deliberative methods to solicit the views of key stakeholders. Despite the concerns that have been raised about data sharing, including consent, ownership of data and privacy issues, there seems to be a growing consensus on the importance of sharing research data more broadly with the scientific community.

64. In this context, Big Data phenomena like real time analysis, linking and sharing of large databases, and use of databases for different purposes in health care and research at the same time and again for different purposes at a later time, make it increasingly problematic and unfeasible to solve the ethical and legal challenges by concepts of ownership. Due to technical equipment, skills and financial resources, only few companies and institutions can deal with very large databases, thus excluding others from gaining knowledge and developing beneficial tools for the health of all. Thus traditional concepts of ownership seem no longer an adequate normative framework.

65. The phenomenon of Big Data rather raises the need for developing new concepts of approaches to balance legitimate interests and benefits. It is necessary to reconcile all the rights and interests which overlap in this field, such as those of the person from whom the data derives, those of the researcher, those of the companies and organizations who use the data, and those of society in general who may benefit from such use. The new context can be seen as a great chance to change our traditional vision and develop new ethical and legal ways for a real scheme of sharing benefits. This shifts the focus from ownership to custodianship and the responsibilities of all stakeholders with benefit sharing being a crucial part. Custodianship means the responsibility for the safety and well-being of someone or something and its synonyms represent some ethical values like care, custody (medical ethics), protection and trust to the guardianship or the safekeeping.

66. Against this background, Big Data can be framed as a common good of humankind. Thus advancements and new opportunities provided by science and technology might help reduce and not deepen the inequalities that prevent many human beings from enjoying the highest attainable standard of health, both at the national and the international level. This is in harmony with Article 2 of the UDBHR which states: "(f) to promote equitable access to medical, scientific and technological developments as well as the greatest possible flow and the rapid sharing of knowledge concerning those developments and the sharing of benefits, with particular attention to the needs of developing countries; and (g) to safeguard and promote the interests of the present and future generations" (UNESCO, 2005).

67. The future will be to create the best data infrastructure available for use by anyone in the world, developing new ways to get economic profits through parallel or complementary services or donations. There is the need to improve solidarity not only from the citizens, allowing them to share their data for the common good, but also for companies and private
actors to share their work for the same end. Some projects from different governments, NGOs and private companies under the common name of Open Data are good examples of the near future, where “the best data infrastructure [could] be made available for use by anyone in the world, a powerful platform for experimentation, discovery, and the creation of new and better ways of living” (Nielsen, 2014).

68. The concepts of responsibility and common good reinforce what the IBC has advocated for in a previous Report on the Principle of the Sharing of Benefits: “[b]y accepting the idea of universal human rights, we acknowledge that these advancements cannot be a privilege. Progress cannot deepen the existing faults of inequality between peoples and countries. At the same time, the report acknowledges that solidarity through participation and not beneficence is the bond of sharing that needs to be boosted. This is the only way to bring together development and respect for all” (UNESCO, 2015). This includes the concepts of capacity building and science education, brain circulation, open access to health-related information, and empowerment and participation in the production of knowledge.

69. Several programmes have been initiated to strengthen capacities of researchers and research institutions across the globe, especially in developing countries. In the context of genomics and biobanking research in Africa, for example, the Human Heredity and Health in Africa Initiative (H3Africa) aims to strengthen genomic and biobanking research expertise on the continent to address health problems in both communicable and non-communicable diseases (H4Africa Consortium, 2014). The expectation is that this initiative will enable African scientists to contribute meaningfully to cutting edge scientific projects and also become leaders in the field of genomics, bioinformatics and data science. Ultimately, the aim is to improve the health of African people and the global community.

70. Developing an ethical framework for benefit sharing in the use of big data raises the question on how it will be applied in different settings. A lot will depend on the context in which the data is being used (in research, health care or life-style-settings), the nature of the benefits, and which individuals or groups have a legitimate interest in the use of the data. This diversity can most comprehensively be addressed by a multi-tiered governance model (see Chapter V). It can entail giving a considerable amount of a data based financial profit for the benefit of the group who substantially contributed their data to make insights and products possible.

IV.4. Justice

IV.4.1. Digital Gap

71. Big data needs tools and providers to access and manage them. The digital gap (called digital divide as well) constitutes one of the major challenges in the process of democratizing information and communication, and thus development. Digital technology is, in fact, a major asset for development and for fighting poverty so that it is one of the objectives of the Millennium Development Goals of the United Nations regarding the strengthening of the global partnership for development. As a result, the gap is diminishing. One target was to ensure that the benefits of new technologies, and especially ICTs, were made available for everyone. This program actually increased the level of information of the populations, in particular the poorest and the most isolated. Regarding the health sector, this strategy strengthened the prevention of health problems thanks to a better health promotion, but also significantly expanded the supply of care through telemedicine in particular. This is a positive development regarding people’s awareness and knowledge, which is the basis of any public health action. The good performances in mobile telephony contributed to the implementation of initiatives in order to strengthen the knowledge and skills for the public and actors in the fight against non-communicable diseases. For example, the Be Healthy Be Mobile program is implemented in several countries (e.g. India, Egypt, Senegal, Costa Rica, Zambia).
72. Internet penetration increased, from just over 6% of the global population in 2000 to 43% in 2015, and 46.4% of the households covered worldwide (ITU, 2015a). Therefore, 3.2 billion people are connected to a global network of apps and contents. However, in many low-income countries, limited international bandwidth and low national infrastructure hinder the provision of broadband Internet services at affordable prices. These limitations have concrete effects on the types of services and applications users can access. Consequently, the developing countries and the least developed countries experiment limitations to integrate the management of Big Data in their offers. In addition, the average price of services remains relatively high in many of the poorest countries where the maintenance and up-dating of the tools (i.e. cell phone) might be difficult.

73. Despite all these advancements in access to ICTs, huge disparities remain and tend to grow between countries, but also within countries. The analysis of Internet household coverage reveals clear differences according to the country’s economic level. Thus, in 2015, Europe had coverage of 81.2%, against coverage of 10.7% for Africa (ITU, 2015b). Barely a third (32%) of the population of developing countries uses the Internet, against 82% in developed countries (ITU, 2015b). Consequently, although problems of isolation and accessibility to health facilities and professionals are given priority in most developing countries, technological innovations such as tele-expertise, tele-diagnosis or tele-consultation, constituting effective solutions are not yet sufficiently implemented.

74. There are also significant inequalities among countries in terms of ICT skills, digital literacy, and the existence of relevant local content. So even if the innovative content is offered through ICTs in developing countries, they cannot be implemented there because it requires infrastructure and human resources for their adaptation and evaluation. This situation raises ethical problems such as massive transfer of data without clear conditions, coming with the risk of “data-piracy”, or inadequate governance policies (Wyber et al., 2015). Indeed, masses of information are transferred from developing countries with limited human and technical resources, to be processed without some assurance on the protection of data transferred.

IV.4.2. Non-discrimination

75. In many countries access to health care is not based on a public model but on private models, which are managed by private insurance companies with varying degrees of involvement, control and monitoring by the public powers. In consequence, subjects can be denied or impeded from accessing insurance due to disclosure of health data. But should insurance companies become aware of the specific risks of a person in detail, it is very difficult to speak of an insurance model which has traditionally been based on balancing of collective risks rather than detailed individual risk profiles.

76. Using ICTs and Big Data approaches, there are coming up behavior-based insurance models in different types of insurance, including health insurance. Persons are offered reduced insurance rates or other kinds of premiums in case they transfer their data e.g. about sportive activities, nutrition habits and the like on a regular basis via tracking, wearables and apps. This can turn out to discriminate against three groups of people either in the form of not having access to benefits or, what is to be expected in the long run, in the form of suffering from disadvantages. Firstly, those people who do not want to share their data with the insurance company are discriminated against, regardless whether they can fulfill the norms like walking 10,000 steps per day or not. Secondly, those people who cannot fulfill the norms because of a disability, a disease or a great misfortune are affected, regardless of their willingness to share the data. But exactly these people are in special need of solidarity and social support. Thirdly, those people who have different beliefs than the insurance companies about what health and a healthy life is about will be discriminated against. The sovereignty to define which health related behavior is rewarded – e.g. walking 10,000 steps but not meditating every morning – should not be with insurance companies. It is rather a scientific and cultural mission.
77. There is also a risk for discrimination coming from Big Data based breaches in anonymization. Aggregating data opens the possibility of re-identification through cross-referencing with data concerning ethnic background, locational data, age, other metadata, health records or genetic data (Choudhury et al., 2014).

78. In addition, there are powerful algorithms, which can profile individuals without having personal data about them in order to form groups according to geographical, socioeconomic, ethnic or other characteristics. The anonymisation of personal data matters little then if the outcomes affect the groups to which they belong with a risk of discrimination and stigmatisation. Such effects impact also those members of the community, who did not give their consent and who didn’t contribute their data but nevertheless are members of the profile-defined group.

79. Furthermore, a loss of freedom can result from a subtle pressure to adjust behavior. Predictions are constructed on the frequency of behavior. The consequence is on the one hand the stigmatization of infrequent behavior (of minorities), and on the other hand a hidden coercive pressure to adopt “frequent behaviour” in order not to be excluded. This pertains to groups as well.

80. There are further possibilities for obvious or subtle stigmatization and structural disadvantages like access to research and progress for those with little equipment and skills in Big Data approaches. Algorithms can contain systematic biases, even ones which eventually contribute to racism. Having this in mind, it has to be assessed with due diligence in which areas and in which ways Big Data can contribute to discrimination. Adequate measures then have to be taken in order to prevent stigmatization and discrimination according to Article 11 of the UDBHR.

IV.5. Sustainability with regard to energy and environment

81. Big Data for health is also an ecological issue both for its energy cost and for the gains it can achieve through a better use of natural resources. Emphasis is put on the potential benefits, including energy savings and the greenhouse problem (Carbon War Room, n.d.). But energy-intensive information technologies are big consumers and produce greenhouse gases at all stages of their life cycle (CNRS, n.d.).

82. There are infrastructures behind a distributed and remote computing that have a high energy consumption and carbon footprint: 10% of world energy, health part unknown but most likely significant. This cost is due for 38% to terminals, 14% to transmission and 48% to data centers (Cappy, 2017). Of note, an estimated 80% of the energy consumption of connected objects (14 billion at present, 125 billion units in 2025) is not dedicated to their use but to maintaining their network connectivity. According to the International Energy Agency the world of connected devices consumed 616 TWh in 2013 (IEA, 2014), which is the consumption of Sweden with its 10 million inhabitants for one year. Data centers bring together thousands of machines running constantly. In addition to the energy required to operate the equipment, the huge needs for cooling must be added. In 2013, US data centers consumed an estimated 91 billion kilowatt-hours of electricity - which would be enough electricity to power all the households in New York City for two years - and are on-track to reach 140 billion kilowatt-hours by 2020 (NRDC, 2015).

83. Improvements might results from voltage equalization, research on quantum computing and bio-inspiration, machine architecture to solve the gap between data management and data storage, changing the voltage in microprocessors.

84. There are also concerns about the possible impact on health by pollution associated with the waste created by the connected but no longer used objects. On a global scale, an estimated 50 million tons of E-waste are produced every year while. The Stopping the E-waste Problem (STEP) initiative (http://www.step-initiative.org/), a joint effort from UN organizations, grassroots groups and industry, predicts that by 2017 the total annual volume of e-waste will have risen by a third, to 65.4 million tons.
85. There are also concerns with the ecological impact due to search and exploitation of essential components to produce around 67 million tons of new electrical and electronic equipment that are put on the market. The term “rare earth” describes a set of 17 chemical elements with exceptional properties. It is thanks to these rare earths that colors of computer screens are so bright, mobile phones have touch screens and wind turbines can generate electricity. The downside is that their extraction and their transformation pollute, produce radioactive waste and distort the landscape. Indonesia and Australia have already been impacted.

IV.6. Research

86. Because Big Data can be a powerful generator of hypothesis, particularly for clinical trials, a primary ethical requisite is to conduct research following the highest standards of ethics, data protection and excellence. Biobanking is at the heart of biomedical research and forms an essential source for Big Data research. Accordingly, best practice guidance concerning Big Data research on biobanked samples and data collected and stored in academic biobanks have been already issued and are regularly discussed through recommendations of international organizations such as BBMRI-ERIC (http://bbmri-eric.eu/) and OECD. Recent advances include committees involving patient representatives reviewing the governance of the biobank including analysis of the use of the potential commercial value of the data, and a process of follow-up information on the various research using the data, allowing the patients to be informed on the long-range and giving them a real possibility to opt-out.

87. In an attempt to confirm the use of highest quality biobanked samples and associated data, the concept of BRIF (Bioresource Research Impact Factor) has been introduced. The BRIF initiative is intended to help in the follow-up of the use of data and databases associated to biological samples. In this context, the CoBRA (Citing of Bioresources in Research Articles) guideline has been developed to guide and standardise citation of bioresources in academic literature. This can be an example for Big Data approaches as well.

88. Understood in terms of “control”, ownership grounds empowerment of data subjects to prevent any kind of data manipulation, including societal pressure to constrain any unacceptable uses (see Chapter IV.3.). For biobanks and databases, control is relevant, e.g. to consider the permissibility of using research data for commercial pursuits as is made possible by allowing private and third party companies to access biobanked samples and data. On the contrary, the ownership of the data may be an obstacle for research, for example preventing certain forms of data-mining or meta-analysis. National laws may limit such research, particularly if a transfer of data is involved.

89. The push towards open data is in contradiction with protection of sensitive medically relevant data. In this context, it is noteworthy that some scientific journals require to transfer all data sets used to raise results to a repository in order to allow other scientists to replicate the results or to use them in meta-analysis. But this may also lead the review to take rights on the data sets and makes data-mining prohibited or too costly. Several national laws are presently under discussion to allow free access to data sets for research.

90. A major challenge is education of citizens to get the best from Big Data derived personal information. Subjects consenting for research must be able to exercise their access rights with reasonable effort so that the right can be exercised meaningfully. Big Data requires advanced scientific knowledge through education. Datasets do not “contain the world in small” and there is a risk that the comprehensive profile becomes a representation of the profiled (Bowker, 2014).

91. A risk exists of an overreliance in Big Data power which has led to the recent development of a direct relationship between data managers and patients/citizens bypassing medical intervention. Such practice does not exist in the academic field but is rapidly
developing in private research by commercial companies. Medical intervention and interpretation are needed because predictions based on big data analysis alone do not take into account factors that require interpretation by medical professionals. But to interpret the meaning of Big Data results for a given person remains a fundamental goal of the relationship between patient and physician.

92. Big Data analyses will reveal unexpected results with potential health relevance for a given individual, so-called incidental findings. Since these questions have been treated in a previous Report of the IBC on the Principle of Non-Discrimination and Non-Stigmatization (UNESCO, 2014), there is no need here to further comment.

93. At the level of public health research, Big Data will be used to develop public health policies to promote health and prevent disease at the aggregate level. This process must not compromise the rights to privacy of individuals and must adhere to good governance on the use of data. The cost of acquiring, maintaining and using Big Data in public health policy is also of major concern, especially for developing countries with scarce resources.

94. Health research now uses connected personal devices such as smartphones for tracking and monitoring to gather health data (mHealth). This can be a helpful source for real world data but it also raises ethical issues such as risk of exploitation or violation of the right to autonomy since mobile phones can collect a wide range and quantity of personal information from their user with little precautional measures to protect autonomy and privacy.

V. GOVERNANCE

95. Governance systems for Big Data should protect the fundamental rights of the persons from whom the data originates, including their freedom to make decisions, and aim at maintaining the trust of the public. Thus, guiding principles for governance has to include autonomy and the right to information, voluntariness, privacy and data protection, transparency, equality and lawfulness. Data governance should guarantee that citizen involvement, engagement, participation and sharing of data will not be subject to exploitation, manipulation and control. The IBC has based its recommendations on these principles.

96. Furthermore individual interests and rights as in public health policies can be balanced against what is called the public good. In the case of Big Data, most of the relevant actors collecting and using data generally lack a democratic mandate from the people whose information is included (willingly or unwillingly) in databases to define what the public good actually is about. This creates an ethical tension that must be dealt with by an appropriate governance structure which could guarantee a “public good”.

97. In view of the global scope of Big Data and health, the huge variety of – ever new – players in this field and the fast technological development it is difficult to elaborate comprehensive and balanced regulations. But in order to make benefits possible, to ensure responsible conduct and to prevent harm prudent governance is required on different tiers and in different ways, guiding and binding the different stakeholders and institutions on the legal level as well as other levels in form of rules, regulations, decrees, agreements, self-binding code of conducts etc.

98. In the era of Big Data, it has become increasingly difficult to rely on the idea that data protection can be regulated by adhering to the paradigm of “consent or anonymize” (see also chapter on autonomy). This leads to the conclusion that it is crucial to develop and apply a comprehensive multi-tiered governance structure, which doesn’t exist yet, for responsible use of data. It is also crucial to note, and has been noted already in paragraph 35, that while in this time and age the individual becomes more and more transparent because we know more and more about that individual, systems tend to become more and more opaque. This needs to be countered by creating more transparent governance systems. Such a governance system can follow different models, but in all cases should be
a public structure, or at least a public-private partnership. Serious participation of patients and the public in setting up and applying the governance is a prerequisite for this.

99. Transparency about assumptions used in the algorithms, which at the moment is mostly opaque, is an essential element in this context. Institutions or companies should be transparent about the algorithms that they use. Systems that use algorithmic decision-making should offer explanations regarding both the procedures followed by the algorithm and the specific decisions that are made. Some even claim that the source code of an algorithm should be published.

100. The fundamental right of data protection offers a conceptual and legal basis for guaranteeing a morally and legally acceptable use of data and for setting up a complex system of information governance rules. The principle of lawfulness, a key principle of data protection, requires that every processing of personal data needs a legitimate legal basis. Its essence is the overarching system of checks and balances, regulating personal data processing activities. It aims to complement the individual rights, guaranteed by the fundamental rights to privacy, with an effective allocation of responsibilities and duties of those who use the data. The principle of lawfulness has its regulatory limits at the different regulations in different countries.

101. The governance framework should address regulatory and ethical issues related to data collection, access, release, and linkage, analysis etc. Against the background of the ethical and legal considerations, the following elements are perceived to be crucial for such a framework. They could be further developed into international treaties, self-binding regulations between providers or may even lead to the creation of a global agency of vigilance. Existing entities such as Research Ethics Committees (RECs), Data Access Committees (DACs) and data protection agencies could play an important role in such multi-tiered governance framework. This is a multi-tiered governance framework, addressing relations between multiple stakeholders.

   a. A clear statement of the purpose of the database.
   b. Procedures for (broad) consent, re-contact (including return of results) and re-consent.
   c. Procedures for dissent, as alternative to consent.
   d. Arrangements for ensuring the rights to access, to rectify, to cancel data.
   e. Arrangements for withdrawal, and a description of the extent to which withdrawal is technically possible.
   f. Arrangements for the protection of privacy, at least declaring the limits of privacy protection.
   g. Policies after the death of a participant.
   h. Arrangements on ownership of the data and the products derived from them.
   i. Arrangements for the collection, storage and duration of storage of data including quality control and safeguards to protect privacy and confidentiality.
   j. Arrangements for access to data including arrangements for data sharing, and criteria for access.
   k. Role of REC or DAC in decision-making about sharing of data.
   l. Arrangements for how the data will be dealt with in the event of change of ownership or closure.
   m. Transparency of the algorithms used for pattern recognition; arrangements to check profiling of individuals or groups according to ethical considerations.
   n. Disclosure of commercial interest and collaboration with commercial parties.
   o. Arrangements that enable participants to remain informed of ongoing and novel use of data including research activities.
p. A clear policy on the disclosure of aggregate and individual research results to participants.
q. Arrangements for the involvement of participants in designing governance procedures, particularly with regard to ethical oversight and communication with data-providers.
r. Arrangements for benefit-sharing.

102. The IBC recommends that an International Legal Instrument on Data Protection is adopted by the Member States. If so requested, the IBC is willing to contribute to the development of a suitable governance framework that should precede such a treaty. Each country can then create within its own jurisdiction country specific legislation, under which an agency can be created that is tasked with the oversight of these governance systems. Such an agency would also provide for a clear point of entry for public control.

VI. RECOMMENDATIONS

103. In order to make Big Data a global success for health, to harvest the opportunities of Big Data in healthcare and research as well as international cooperation, while at the same time avoiding violation of fundamental human rights stipulated in the Universal Declaration of Human Rights and in the Universal Declaration on Bioethics and Human Rights, the IBC gives the following recommendations.

104. Facing the complex nature, the global scope, and the wide variety of stakeholders involved in Big Data related to health, a multi-tiered governance approach is crucial, so that trust and control are balanced for the benefit of all (for more detail see Chapter V).

105. The IBC considers four measures to be crucial for protecting individual rights and fostering public good while recognizing the unavoidable loss of control by individuals about the use of their data in times of Big Data: governance, education, capacity building, and benefit sharing.

106. In the following, the IBC presents examples of important measures. They are understood as a step in an ongoing global debate. They address different stakeholders, but finally the success of these measures will depend on comprehensive cooperation and participation of all stakeholders including patients, research subjects and citizens in general.

107. International Agencies are called upon to develop and support a global framework for the use of Big Data in health related areas, especially in healthcare and research. For example:

a. The UN is recommended to develop and adopt an International Legal Instrument on Data Protection.

b. UNESCO is recommended to develop a convention on the protection of privacy, including a framework for new approaches to ownership and custodianship of personal data.

c. WHO is recommended to conclude an agreement with app-stores about presenting health-related apps in a way that autonomy, transparency and adequate information are guaranteed.

d. A global vigilance system and a global agency of vigilance for the use of Big Data in health related applications should be set up.

e. International Agreements should adopt an understanding of Big Data as a common good of humankind and facilitate open access and use of Big Data for the common good where feasible. For this purpose a public data infrastructure should be set up.
f. OECD is recommended to develop a framework for sharing of benefits from Big Data applications. This can also contribute to increasingly overcome the digital gap.

108. National Governments are called upon to develop and launch an action plan including legislation and policies which should among others address the following aspects:

   a. Internationally harmonized implementation of globally accepted data protection principles.
   b. Capacity-building for Big Data-related healthcare and research including an efficient data infrastructure.
   c. Promotion of learning healthcare systems.
   d. Enabling effective cross-border cooperation in the processing of personal health data for health-related public interest purposes.
   e. Promotion of education with regard to Big Data-related skills, competences, and awareness about ethically relevant implications. Particular attention should be paid to vulnerable groups like minors and people with impaired capacities.
   f. Context specific diversified models of consent should be introduced and specified, allowing for broad and dynamic consent where appropriate.
   g. Privacy of individuals sharing their data should be protected by default in every processing step as well as by design of the device and technological equipment. Separate attention should be paid to the protection of group privacy – not least against the background of the manifold possibilities of discrimination.
   h. An Research Ethics Committee or a similar institution should be implemented to oversee commercial Big Data research.
   i. Instruments for meaningful public and patient involvement should be developed and widely implemented in a culture sensitive way.

109. The wide variety of healthcare and research institutions as well as companies, informatics developers and other stakeholders are called upon to develop context sensitive guidelines, code of conducts and self-binding instruments with a special focus on Big Data applications. Ethical policies and codes of conduct should be developed for health professionals, computer scientists, clinical researchers, data scientists and other stakeholders of an institution. Motivating ethical behavior should be the primary goal, but consequences of violation should also be provided for.
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