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SAFEGUARDING THE GENOME: HARMONIZING PRIVACY AND PROGRESS IN INDIA'S STRUGGLE AGAINST GENETIC DISCRIMINATION

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ABSTRACT

Genetic testing opens up amazing chances for spotting diseases early and tailoring healthcare, but it also brings real risks when personal genetic info is misused. Discrimination based on genetic data can lead to missed job opportunities, higher insurance premiums, and social stigma, which threatens individual rights and public trust. This article looks at India's messy legal framework—rooted in broad constitutional promises of equality and privacy, along with various scattered laws and guidelines—and compares it to more coherent international models like the U.S. Genetic Information Nondiscrimination Act (GINA), HIPAA, and the EU's GDPR. By doing a critical side-by-side analysis, it highlights key shortcomings in India's strategy, such as the lack of a specific anti-discrimination law, weak ethical codes, and enforcement issues. In the end, the article suggests a practical reform plan: creating a standalone Genetic Non-Discrimination Act, updating data-protection laws to treat genetic data as extra sensitive, setting up an independent Genetic Data Regulatory Authority, and rolling out focused awareness and training programs. These steps aim to strike a good balance between privacy and progress, ensuring that India can embrace genetic advancements responsibly while respecting individual dignity and fairness.

Keywords: Genetic Discrimination; Privacy Law; India; Employment and Insurance; Legal Reform

(I) INTRODUCTION

Genetic testing is making huge strides and changing the way we approach healthcare. With the ability to catch diseases early, create personalized treatment plans, and pinpoint hereditary risks, these genetic technologies are set to transform medical care and preventive health. Just a simple test can provide people with insights about their chances of facing conditions like cancer, diabetes, or heart disease, which can lead to timely actions that might save lives. But as we dive into these exciting advancements, some serious issues are popping up regarding the potential misuse

of genetic data—especially when personal information leaks out of the doctor's office.

One of the biggest concerns revolves around jobs and insurance. If genetic information is accessed or interpreted incorrectly, it could lead to missed job opportunities, impact hiring decisions, or even result in higher insurance premiums for individuals based on their predicted health risks. Picture a scenario where someone, genetically likely to develop a neurological issue, gets quietly pushed aside in a job opening, or a young person faces sky-high life insurance rates after a test hints at a future cancer risk. These situations aren't just

hypothetical; they're already happening in various places, raising serious questions about fairness, privacy, and respect for individuals.

Given this context, it's crucial to look into the laws that are supposed to protect people from genetic discrimination. While some countries have solid protections in place, India's approach currently feels all over the place and isn't doing enough, leaving many people vulnerable. This article aims to take a closer look at India's legal landscape regarding genetic privacy and discrimination, compare it to international standards, and suggest a practical way forward. The goal is to join the important national discussion on how India can make the most of genetic technologies while still protecting everyone's fundamental rights.

(II) UNDERSTANDING GENETIC DISCRIMINATION AND ITS ETHICAL BASIS

Genetic discrimination is all about treating people unfairly based on genetic info that shows they might be prone to certain diseases or traits. As genetic testing gets easier and more accurate, worries have grown about how it's used in job screenings and insurance decisions, where these test results could lead to people getting excluded, stigmatized, or punished because of their DNA. When we look at the ethical angles, three big ideas come into play: privacy—making sure folks control who gets to see and use their info; autonomy—ensuring that testing is done with informed consent and is totally voluntary; and equity—protecting those who are more vulnerable from getting hit harder. It also brings up some tough moral questions about fairness and justice in society.

Direct vs. Indirect Genetic Discrimination

Direct genetic discrimination happens when choices about someone are plainly based on their genetic info. For instance, an employer might not hire, promote, or keep a worker just because they have a genetic tendency toward something like cancer or a neurological issue. Likewise, an insurance company could refuse

coverage or slap on higher rates if they see genetic test results, even if the person is perfectly healthy at the moment. This type of discrimination is pretty obvious and can lead to serious problems like job loss, emotional stress, money troubles, and social stigma.³¹⁶

Indirect genetic discrimination is sneakier and often hides within policies or practices that seem neutral but actually hurt people based on their genetic traits. For example, a company's wellness program might offer cash rewards only to employees who get genetic screenings, which indirectly punishes those who choose not to test because they're worried about their results being misused. This form of discrimination is trickier to spot and deal with since it usually links with broader systemic issues and personal choices.³¹⁷

Ethical Foundations of Genetic Discrimination

Genetic privacy is a key ethical principle that protects people from having their DNA info collected, used, or shared without permission. With the rise of genetic testing technology, worries have grown about how this sensitive data is handled. Things like employers and insurers can easily access someone's genetic profile, often without their knowledge. When genetic privacy is breached, it doesn't just lead to discrimination; it can also cause emotional distress, social stigma, and a breakdown of trust in healthcare and genetic research.³¹⁸

Genetic autonomy is all about an individual's right to make informed choices about their own genetic data. This means they get to decide whether to take genetic tests, if and when to share their results, and how their genetic information is used. Respecting genetic autonomy is essential to avoid situations where people feel pressured to disclose sensitive information at work or with insurance

³¹⁶ Akash Akash and Shuvro Prosun Sarker, "Analysing Discrimination Based on Genetic Information" 11 *Lentera Hukum* 157 (2024).

³¹⁷ N. V. Chernyh, "Genetic and Digital Discrimination in Labor Law: Problem Statement" *Courier of Kutafin Moscow State Law University (MSAL)* 48-57 (2024).

³¹⁸ Anya Prince, "The Genetic Information Privacy Act" 330 *Jama* 2049 (2023).

companies because they fear negative consequences

Genetic equity aims to ensure that genetic information isn't used to reinforce or create new types of social, racial, or economic inequalities. It advocates for fairness and works to prevent discrimination, especially against marginalized groups that may be more affected by certain genetic traits.

Effects on Different Demographics

At-risk groups, like racial and ethnic minorities, might face a bigger impact from genetic discrimination. These communities often deal with existing barriers in healthcare and jobs, and misusing genetic info could make things even worse for them. It's super important to ensure fairness in genetics to protect these groups from getting hurt further. People who have known genetic disorders or are at risk for them can experience discrimination when it comes to work and insurance. For instance, someone with a genetic likelihood of developing a chronic illness might get turned down for life insurance or have to pay higher rates, even if they're totally healthy. This kind of discrimination can lead to money troubles and make it harder for them to get the services they need. The impacts of genetic discrimination reach beyond individual issues. If this becomes common, it might make people think twice about getting genetic tests, which could slow down progress in genetic research and personalized medicine. Plus, it could shake public confidence in institutions, leading to social tensions and even more inequality.³¹⁹

(III) INTERNATIONAL LEGAL STANDARDS

The legal framework regarding genetic privacy has undergone significant transformations, largely influenced by advancements in genomic technology. Nations across the globe are working to reconcile individual rights to genetic privacy with the collective interests in

research and healthcare, thereby formulating legal standards to combat discrimination and safeguard sensitive genetic information.

International legal frameworks

These are pivotal in the development of genetic privacy laws. These frameworks often serve as benchmarks for national legislation, highlighting the necessity of safeguarding genetic data while promoting scientific advancement.

The General Data Protection Regulation (GDPR), implemented by the European Union, is considered the benchmark for keeping data safe. It treats genetic data as a special type of personal information that needs extra protection. The GDPR focuses on getting consent, limiting data collection, and being transparent, and it also lays out legal grounds for using genetic data in research and public health.³²⁰

The Universal Declaration on the Human Genome and Human Rights, adopted by UNESCO in 1997, lays out ethical principles for the safeguarding of human genetic data. It stresses the prohibition of genetic discrimination and emphasizes the significance of informed consent. Another instrument by UNESCO, the International Declaration on Human Genetic Data, offers guidelines for the collection, processing, and dissemination of genetic data. It highlights the importance of international collaboration and the protection of vulnerable populations.³²¹ These international frameworks underscore the necessity of striking a balance between individual rights and societal advantages, ensuring that genetic data is utilized in a responsible and ethical manner.

Regional Legal Frameworks:-

European Union (EU)
The European Union (EU) has established a comprehensive and robust legal framework

³¹⁹ Alexander Nill, Gene R. Laczniak, *et.al.*, "The Use of Genetic Testing Information in the Insurance Industry: An Ethical and Societal Analysis of Public Policy Options" 156 *Journal of Business Ethics* 105-121 (2017).

³²⁰ Vishal Kumar Seshagirirao Anil and Adeoluwa Bennard Babatope, "Data Privacy, Security, and Governance: A Global Comparative Analysis of Regulatory Compliance and Technological Innovation" 21 *Global Journal of Engineering and Technology Advances* 190-202 (2024).

³²¹ Lyudmila N. Berg, "Genomic Data Privacy and the Right to Science: Issues of Balance" 77 *Lex Russia* 93-101 (2024).

aimed at safeguarding genetic privacy, primarily through the General Data Protection Regulation (GDPR). According to the GDPR, individuals are required to provide explicit and informed consent prior to the processing of their genetic data and maintain the right to withdraw such consent at any point. Organizations are mandated to implement data protection principles via "Data Protection by Design and Default," ensuring that protective measures are incorporated from the outset. Furthermore, the GDPR imposes strict regulations concerning cross-border transfers of genetic data, necessitating adherence to strong protections or compliance with EU privacy standards.³²²

United States (US)

In the United States, the legal landscape surrounding genetic privacy is characterized by its fragmentation, comprising an amalgamation of both federal and state regulations. The Health Insurance Portability and Accountability Act (HIPAA) safeguards genetic information in healthcare environments; however, its coverage is not comprehensive across all scenarios involving genetic data utilization. Furthermore, the Genetic Information Nondiscrimination Act (GINA) explicitly prohibits the utilization of genetic information for discriminatory practices related to employment and health insurance, yet it notably falls short of extending protections to domains such as life insurance or disability insurance. Additionally, the California Consumer Privacy Act (CCPA) establishes substantial safeguards for genetic information, thereby setting significant precedents for other states to consider in their regulatory frameworks.³²³

Asia-Pacific region

Countries within the Asia-Pacific region have implemented diverse legal strategies to tackle concerns related to genetic privacy. In Japan, the Act on the Protection of Personal

Information (APPI) categorizes genetic data as sensitive personal information, requiring stringent measures to safeguard its confidentiality. Meanwhile, South Korea's Personal Information Protection Act (PIPA) places a strong emphasis on transparency regarding the utilization of personal data and advocates for the localization of data processing to uphold citizens' privacy rights.³²⁴ Conversely, India's Criminal Procedure (Identification) Act, 2022³²⁵, allows for the collection and retention of genetic data for criminal investigations, prompting significant discussions surrounding the balance of proportionality and the adequacy of privacy protections.

IV) INDIA'S LEGAL SYSTEM: STATUTES AND CONSTITUTIONAL RIGHTS

Constitutional Rights: Articles 14 and 21

The Constitution of India enshrines fundamental rights, providing a crucial basis for safeguarding genetic privacy and curbing discrimination, even though these rights were not specifically formulated with genetic science in view. Notably, Articles 14 (equality before the law) and 21 (right to life and personal liberty) can be interpreted to encompass genetic issues.

Article 14: Equality before the Law ensures that "the State shall not deny to any person equality before the law or the equal protection of the laws." Historically, this provision has tackled discrimination based on religion, race, caste, sex, or place of birth. However, its expansive language allows for the extension to newer forms of discrimination, including those rooted in genetic characteristics. Within this framework, any legislation or policy that differentiates between individuals based on genetic predispositions—such as denying employment to an individual with a BRCA mutation—can be contested as contravening Article 14's guarantee of substantive equality.³²⁶ In practice,

³²² Pieter De Smet and Mahsa Shabani, "Genetic Data Sharing in the View of the EU General Data Protection Regulation (GDPR)" *Genomic Data Sharing* 171-187 (2023).

³²³ *Supra* (note 5)

³²⁴ *ibid*

³²⁵ Criminal Procedure (Identification) Act, 2022

³²⁶ *Supra* (note 1)

litigants could assert that genetic discrimination engenders an “arbitrary classification” devoid of a reasonable connection to a legitimate state objective, thereby invoking Article 14’s stringent scrutiny.

Article 21: Right to Life and Personal Liberty upholds “the right to life and personal liberty,” a provision the Supreme Court has broadly interpreted to encompass personal privacy. In *K.S. Puttaswamy v. Union of India* (2017), the Court unanimously affirmed that informational privacy is essential to individual dignity and autonomy, explicitly acknowledging the right to govern personal data, including health and genetic information).³²⁷ This landmark ruling establishes a constitutional basis for the argument that unauthorized collection, dissemination, or exploitation of genetic data constitutes a violation of Article 21.

Limitations of Articles 14 and 21

Despite these constitutional safeguards, Articles 14 and 21 do not explicitly confront the distinctive challenges posed by genetic data—namely, its predictive capabilities, familial ramifications, and the potential for stigmatization. Article 14 does not recognize genetic traits as a forbidden ground for discrimination, and Article 21 does not specify the procedural protections necessary for managing sensitive genomic data. In the absence of specific legislation, such as a Genetic Non-Discrimination Act, individuals must rely on these broad protections, which do not feature specialized enforcement mechanisms or distinct remedial provisions for genetic discrimination.³²⁸ Therefore, while Articles 14 and 21 present viable pathways for judicial engagement, there remains an urgent necessity for comprehensive statutory measures to address these significant deficiencies.

³²⁷ Nitin Raj Singh, “Right to Privacy in India: Constitutional Basis and Implications on Modern Issues” III *The Journal of Unique Laws and Students* (2024).

³²⁸ *Supra* (note1)

Statutes and Guidelines:-

India's framework for protecting genetic data is characterized by a patchwork of statutes and policy guidelines, which, while recognizing the sensitive nature of genetic information, lack robust anti-discrimination protections.

Information Technology Act, 2000

The Information Technology Act provides the legal structure for electronic data privacy and security. According to Section 43A, any corporate entity handling “sensitive personal data or information” (SPDI) is required to adopt “reasonable security practices” to avoid civil liability for negligence. The accompanying SPDI Rules (2011) classify medical records and history as SPDI, implicitly including genetic data, thus obliging companies to secure such information and obtain consent before collection or dissemination. However, the absence of explicit references to genetic profiles within the Act generates ambiguity regarding the pursuit of genetic discrimination claims under its provisions.³²⁹ Enforcement is additionally hindered by inadequate public awareness of the Act’s SPDI stipulations and procedural barriers to compensation via adjudicating officers.

Pre-Natal Diagnostic Techniques (PNDT) Act, 1994

Designed to prevent sex-selective abortion, the PNDT Act regulates prenatal diagnostic methods, including genetic tests for identifying chromosomal anomalies. It necessitates strict confidentiality concerning test results and imposes penalties for unauthorized disclosures.³³⁰ Although this confidentiality requirement indirectly supports genetic privacy, the Act’s narrow focus on combating female feticide fails to address broader concerns surrounding genetic discrimination. There exists no framework within the PNDT Act to contest the utilization of genetic data for non-medical

³²⁹ Vidhi Agarwal, “Privacy and Data Protection Laws in India” 5 *International Journal of Liability and Scientific Enquiry* 205 (2012).

³³⁰ Lily Srivastava, “DNA Legislations, Challenges, and Opportunities” *Handbook of DNA Forensic Applications and Interpretation* 119-147 (2022).

purposes such as employment or insurance underwriting.

ICMR Bio-banking Guidelines

The Indian Council of Medical Research's 2017 National Ethical Guidelines for Biomedical and Health Research delineate best practices for bio banks, mandating informed consent, anonymization of samples, and oversight by institutional ethics committees. These guidelines emphasize secure storage, restricted access, and explicit data-sharing agreements. Nevertheless, as these guidelines are nonbinding, adherence relies on the goodwill of research institutions and the vigilance of ethics committees, offering limited recourse for individuals in the event of misuse of their genetic samples.³³¹

Criminal Procedure (Identification) Act, 2022

The Criminal Procedure (Identification) Act permits law enforcement to gather and retain biometric identifiers, including genetic samples, for the investigation of designated offenses. While this enhances forensic capabilities, the Act does not confine the utilization of such data to criminal justice pursuits; it also lacks provisions for the destruction of samples post-case resolution or independent oversight. The absence of clear procedural safeguards intensifies the potential for genetic information, collected for criminal investigations, to be repurposed by other governmental agencies or private entities, raising significant privacy concerns.

Overall, these statutes and guidelines demonstrate India's acknowledgment of the sensitive nature of genetic data while exposing critical shortcomings: the lack of explicit anti-discrimination measures, reliance on nonbinding guidelines, and inadequate enforcement mechanisms. Therefore, comprehensive legislative reform is imperative to address these discrepancies and safeguard individuals from genetic discrimination across all sectors.

Gaps in Enforcement:-

Nonbinding

The guidelines for bio banking established by the Indian Council of Medical Research (ICMR) are categorized as nonbinding, which indicates that they do not possess legal authority. Consequently, this status leads to disparities in their implementation and enforcement within various research institutions and bio banks. In the absence of legal support, these guidelines are unable to guarantee consistent standards for data protection and privacy.³³²

Low

In India, the public's understanding of genetic privacy and the associated risks of genetic data remains insufficient. This limited awareness impairs the effective application of existing laws and guidelines, as individuals may not fully grasp their rights or the potential ramifications of disclosing their genetic information.

Absence of Standalone Anti-Discrimination Law

India does not possess a distinct legal framework targeting genetic discrimination. Although Articles 14 and 21 offer certain protections against discrimination, they do not specifically address the unique challenges presented by genetic information. This deficiency renders individuals susceptible to discrimination based on their genetic characteristics.³³³

(VI) COMPARATIVE ANALYSIS OF GENETIC DISCRIMINATION METHODS: INTERNATIONAL LAWS VS. INDIA

Legal India's

India currently does not possess a dedicated Genetic Non-Discrimination Act, instead relying on overarching constitutional guarantees and penal provisions to address matters related to genetic discrimination. While the Indian legal framework provides protections under Articles 14 and 21 of the Constitution, which affirm the

Codes

Awareness

Frameworks:- Approach

³³¹ Sachin Chaturvedi, Krishna Ravi Srinivas, *et.al.*, "Biobanking and Privacy in India" 44 *Journal of Law Medicine & Ethics* 45-57 (2016).

³³² *Supra* (note 16)

³³³ *Supra* (note 1)

rights to equality and privacy, these provisions are general and fail to specifically address the nuances associated with genetic information. Consequently, individuals in India are at considerable risk of abuse of genetic data, particularly in sectors such as employment and insurance, where the potential for discrimination is notably pronounced.³³⁴

The lack of targeted legislation on genetic discrimination puts India at odds with advancements in other jurisdictions. For instance, the United States has enacted the Genetic Information Nondiscrimination Act (GINA)³³⁵, which specifically prohibits employers and health insurers from utilizing genetic information in decisions regarding hiring, termination, coverage, or premium rates. Similarly, Canada's Genetic Non-Discrimination Act³³⁶ (GNDA) criminalizes the requirement for individuals to undergo genetic testing or disclose genetic test results as a condition for receiving goods, services, or employment opportunities. These explicit legislative frameworks serve as robust examples of genetic rights protection, a domain in which India currently falls short.

International Laws

Globally, numerous nations have taken proactive steps to legislate against genetic discrimination. Within the European Union, Article 21 of the EU Charter of Fundamental Rights specifically prohibits discrimination based on genetic characteristics. However, the implementation of these principles varies significantly across member states. Countries such as France and Germany have established comprehensive laws governing the usage of genetic information, while others still rely on general human rights protections.³³⁷

In contrast to India, many international frameworks emphasize the necessity for binding, sector-wide regulations to

comprehensively address genetic discrimination. Experts widely concur that explicit prohibitions within anti-discrimination legislation are essential for recognizing and safeguarding genetic rights as a fundamental human right.³³⁸

Public Awareness and Education:- Challenges in India

The level of public awareness regarding genetic discrimination in India remains low, accompanied by insufficient education concerning the risks and ramifications associated with genetic testing. This deficiency in awareness heightens individuals' susceptibility to genetic discrimination, as they may lack knowledge of their rights and the protections accessible to them. The failure to implement a coordinated public education initiative further exacerbates this issue, leaving many individuals uninformed about the potential dangers of genetic testing and the potential misuse of genetic information.³³⁹

International Comparisons

In the United States, the general understanding of genetic discrimination legislation, such as the Genetic Information Nondiscrimination Act (GINA), is notably limited. Research indicates that a significant portion of respondents are not aware of any legal protections against genetic discrimination, with many mistakenly believing that GINA provides more comprehensive safeguards than it actually does.³⁴⁰ Similarly, in Canada, the rollout of the Genetic Non-Discrimination Act (GNDA) has encountered uncertainty and confusion both within the public sphere and the medical community, emphasizing the necessity for enhanced public education and awareness initiatives.

³³⁸ Yann Joly, Charles Dupras, *et al.*, "Looking Beyond GINA: Policy Approaches to Address Genetic Discrimination." 21 *Annual Review of Genomics and Human Genetics* 491-507 (2020).

³³⁹ Hannah Kim, Calvin W. L. Ho, *et al.*, "Genetic Discrimination: Introducing the Asian Perspective to the Debate." 6 *Npj Genomic Medicine* (2021).

³⁴⁰ Lindsay Willard, Wendy R. Uhlmann, *et al.*, "The Genetic Information Nondiscrimination Act and Workplace Genetic Testing: Knowledge and Perceptions of Employed Adults in the United States" 34 *Journal of Genetic Counseling* (2024).

³³⁴ *ibid*

³³⁵ Genetic Information Nondiscrimination Act, 2008

³³⁶ Genetic Non-Discrimination Act, 2017

³³⁷ Aisling De Paor and Delia Ferri, "Regulating Genetic Discrimination in the European Union" 17 *European Journal of Law Reform* 14-32 (2015).

The European Union has also encountered similar obstacles in its efforts to improve public awareness of genetic discrimination. Despite the establishment of legal frameworks, a significant number of individuals remain unaware of their rights and the protections provided by EU regulations.³⁴¹ This widespread lack of awareness highlights the urgent need for comprehensive public education programs to ensure that individuals are well-informed about the risks and implications of genetic testing as well as the legal safeguards available to them.

Technological Trends and Their Impact:- Direct-to-Consumer genetic testing

The emergence of direct-to-consumer (DTC) genetic testing has posed new challenges both in India and on an international scale. DTC testing enables individuals to obtain their genetic information independently of healthcare providers, which raises significant concerns regarding the potential misuse of genetic data by employers and insurance companies. In India, the absence of specific legislation governing DTC testing has resulted in a regulatory void, thereby exposing individuals to the risk of genetic discrimination.³⁴²

On a global level, DTC testing has similarly outpaced the development of regulatory frameworks, with numerous countries facing difficulties in keeping abreast of the rapid growth within the genetic testing market. For instance, in the United States, DTC testing has prompted concerns regarding the sufficiency of the Genetic Information Nondiscrimination Act (GINA), as the statute does not explicitly prohibit the use of genetic information in contexts such as life insurance.³⁴³

Gene-Editing and Emerging Technologies

The regulatory complexities have been further intensified by emerging technologies, particularly gene-editing tools like CRISPR. While these innovations hold the promise of transforming healthcare, they concurrently

introduce ethical and legal dilemmas associated with the potential misuse of genetic data.³⁴⁴ In India, the absence of dedicated legislation for these technologies leads to a deficiency in the legal framework, thereby rendering individuals susceptible to exploitation.

Globally, the oversight of gene-editing technologies remains a pressing issue. Many nations are grappling with the challenge of formulating comprehensive regulatory frameworks that appropriately weigh the advantages of these technologies against the risks of genetic discrimination. Although the European Union has initiated measures to tackle these challenges, the lack of uniformity among member states' laws results in inconsistencies in protective measures.³⁴⁵

Trust in Healthcare and Labor Markets:- India's Trust Gap

The absence of specific legislation in India has significantly diminished trust in both the healthcare and labor sectors. Individuals may be hesitant to pursue genetic testing or disclose their genetic information due to apprehensions regarding discrimination. This prevailing mistrust can have profound implications for public health, as individuals might forgo genetic testing, which could yield crucial insights into their health risks.

International Trust Issues

On an international scale, concerns about trust in healthcare and labor markets persist. In the United States, insufficient public awareness regarding the protections afforded by GINA has perpetuated anxieties about genetic discrimination, thereby dissuading individuals from engaging in genetic testing or research. Likewise, in the European Union, the lack of cohesive legal protections has instigated uncertainty and skepticism among the populace.³⁴⁶

The prevailing distrust in healthcare and labor markets highlights the urgent need for

³⁴¹ Supra (note 22)

³⁴² Supra (note 23)

³⁴³ Supra (note 25)

³⁴⁴ Supra (note 23)

³⁴⁵ Supra (note 22)

³⁴⁶ *ibid*

comprehensive legal frameworks and public education programs. In the absence of strong protections and a well-informed public, the potential advantages of genetic research and technology may be compromised by fears of discrimination and exploitation.

(VII) CONCLUSION

The quick rise of genetic technologies has kicked off a major change in healthcare, research, and society overall. However, in India, the lack of a solid legal framework around genetic data puts individuals at risk of discrimination, privacy breaches, and exploitation. While existing constitutional rights to equality and privacy are important, they fall short when it comes to the complex issues created by the misuse of genetic information.

At this crucial point, India needs to go beyond just patching up protections and adopt a strong, proactive strategy. Safeguarding people's dignity, autonomy, and security in this genomic age isn't just a legal requirement—it's a social necessity. If action isn't taken soon, it could erode public trust in science, healthcare, and job markets, hindering the very innovation that India aims to promote.

By learning from the best practices around the world and customizing them to fit India's unique situation, we can build a future where both genetic advancements and human rights can flourish. This future will allow individuals to benefit from genetic technologies without fear, knowing that the law is there to protect them.

Recommendations & Way Forward

1. Make a Comprehensive Genetic Non-Discrimination Act

India should focus on creating and enforcing a specific Genetic Non-Discrimination Act. This Act needs to: Clearly define "genetic information" to address both inherited and acquired traits. Apply across all sectors, including jobs, health insurance, life insurance, education, and public services. Ban any form of genetic discrimination, both

direct and indirect. Ensure strong civil remedies like compensation and penalties for those who don't comply. Draw inspiration from successful policies like the U.S. Genetic Information Nondiscrimination Act (GINA) and Canada's Genetic Non-Discrimination Act (GNDA), while adapting these ideas to fit India.

2. Update India's Data Protection Framework

India's evolving data protection laws need to specifically recognize genetic data as a unique category of "sensitive personal data." Essential obligations should include: Requiring informed consent before gathering or using genetic information. Only collecting the necessary genetic data for a lawful purpose. Conducting privacy impact assessments before implementing large-scale genetic data initiatives. Imposing strict penalties for unauthorized sharing or misuse of genetic data.

3. Establish a Genetic Data Regulatory Authority

There's an urgent need for an independent Genetic Data Regulatory Authority to: Manage the ethical collection, storage, and use of genetic data. Look into and resolve complaints about genetic discrimination. Set clear standards for data security, consent processes, and the secondary use of genetic information. Work with other regulatory bodies to provide comprehensive protection. This authority should include experts from genetics, law, bioethics, public health, and data science to ensure a well-rounded approach.

4. Run Nationwide Public Awareness Campaigns

To make sure laws are more than just words, India needs to put a lot of effort

into educating the public about genetic rights. This should include: Specific campaigns that explain people's rights related to genetic privacy and discrimination. Resources that are accessible to various communities, especially rural and underrepresented groups. Efforts aimed at vulnerable populations at a higher risk of genetic discrimination. Public understanding is key—not just for individual empowerment but also to create societal pressure for compliance.

5. Provide Specialized Training for Employers, Insurers, and Healthcare Providers

Having laws isn't enough to stop genetic discrimination unless those who handle genetic data receive proper training. Employers need workshops on legal hiring practices that respect genetic privacy, insurers should be educated on fair underwriting that doesn't depend on genetic predispositions, and healthcare providers need to be trained on keeping patient confidentiality and autonomy intact during genetic testing. Training must focus on ethical standards, liability risks, and the broader importance of genetic equality.

6. Future-Proof Legislation Against Technological Trends

The law should stay ahead of emerging trends like direct-to-consumer genetic testing, AI-driven genetic risk assessments, and CRISPR gene editing. Future-proofing strategies might include: Covering consumer genetic testing firms under the same protections. Monitoring how genetic data is used secondary to its original purpose (like for marketing or insurance profiling). Keeping tabs on tech advancements for regular updates to the legislation.

7. Balance Innovation with Rights

Lastly, reforms should carefully balance

protecting individual rights while promoting research and innovation. There can be allowances for anonymized genetic research that benefits public health but without compromising individual consent and privacy protections. This balanced approach will help ensure that India stands out as a leader not just in genomic innovation but also in genomic fairness.