

Genetic Ancestry Analysis Reveals Misclassification of Commonly Used Cancer Cell Lines



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Abstract

Background: Given the scarcity of cell lines from under-represented populations, it is imperative that genetic ancestry for these cell lines is characterized. Consequences of cell line mischaracterization include squandered resources and publication retractions.

Methods: We calculated genetic ancestry proportions for 15 cell lines to assess the accuracy of previous race/ethnicity classification and determine previously unknown estimates. DNA was extracted from cell lines and genotyped for ancestry informative markers representing West African (WA), Native American (NA), and European (EUR) ancestry.

Results: Of the cell lines tested, all previously classified as White/Caucasian were accurately described with mean EUR ancestry proportions of 97%. Cell lines previously classified as Black/African American were not always accurately described. For instance, the 22Rv1 prostate cancer cell line was recently found to carry mixed genetic ancestry using a much smaller

panel of markers. However, our more comprehensive analysis determined the 22Rv1 cell line carries 99% EUR ancestry. Most notably, the E006AA-hT prostate cancer cell line, classified as African American, was found to carry 92% EUR ancestry. We also determined the MDA-MB-468 breast cancer cell line carries 23% NA ancestry, suggesting possible Afro-Hispanic/Latina ancestry.

Conclusions: Our results suggest predominantly EUR ancestry for the White/Caucasian-designated cell lines, yet high variance in ancestry for the Black/African American-designated cell lines. In addition, we revealed an extreme misclassification of the E006AA-hT cell line.

Impact: Genetic ancestry estimates offer more sophisticated characterization leading to better contextualization of findings. Ancestry estimates should be provided for all cell lines to avoid erroneous conclusions in disparities literature.

Introduction

In cancer research, biological model systems are used to understand the contribution of cellular mechanisms to tissue function, disease pathogenesis, and drug efficacy (1–4). Human cell lines representing various tissues in normal and disease states are vital tools necessary to establish preclinical research models. Effects and mechanisms of genetic, epigenetic, and chemical perturbations on cellular viability are explored *in vitro* and *in vivo* through gene knockdown, knockout, and transfection assays using human

cell lines (4–7). Cellular assays employing human cell lines have proven essential to determine the impact of intra- and intergenetic polymorphisms on gene expression and cellular viability (8, 9). Although impactful mutations altering gene expression, cell function, or cell viability have been observed, there is significant variability in reproducibility among varying cell lines (10, 11). A portion of this variability is attributable to differences in genetic background (e.g., varying polymorphisms, epigenetic signatures, and gene expression patterns) among cell lines of the same ethnicity as well as between cell lines representing different ethnicities (12–16).

The examination of experimental outcomes using a racial/ethnic variety of cell lines exposed to the same conditions is a useful tool to elucidate genes or pathways driving the biological contribution to disease disparities (15, 17–19). Because genetic tapestries differ within and between populations, it is important to include a variety of cell lines derived from individuals within and between populations in an effort to provide an accurate representation of the genetic variability (20–22). Currently, there are many commercially available cell lines representing various tissues and disease stage. However, the majority of these cell lines are from patients of European descent, and there remains a critical need to diversify the selection (23, 24). As an example of the existing disparity in cellular model diversity, a recent search of the American Type Culture Collection (ATCC) website for cell lines derived from normal and malignant breast tissue revealed 59 specimens designated as Caucasian/White. Conversely, there were only 14 non-Caucasian/White cell lines consisting of 11

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designated as Black, 1 designated as Hispanic, and 1 designated as East Indian (24). Of note, 10 additional breast tissue cell lines lacked any ethnicity assignment (24).

The lack of adequate genetic population representation among human cell lines leads to a cascade of downstream consequences. It is reasonable to assume that numerous variants, genes, and pathways involved in disease pathogenesis or drug efficacy remain undiscovered when the majority of cell lines used for research are derived from patients of European descent. Because there are lower levels of variation within the European population, we miss potential genetic indicators that steer molecular mechanisms, disease outcomes, and drug response (22, 25–27). Also, SNPs identified in European populations do not completely capture the genetic variation contributing to a phenotype in other racial groups (28). More disturbingly, entire demographics are ignored in key mechanistic studies hindered by the restraints monoethnic cellular models afford (29, 30). This is problematic as underrepresented racial/ethnic groups, who are more likely to suffer disproportionate disease incidence and mortality, remain also underrepresented in biospecimens available for research (31–34).

Complementary to the aspiration of health disparity researchers to diversify commercially available biospecimens, the NIH has currently established guidelines concerning the authentication of key biological resources to ensure identity and validity (35). In response to this recommendation, a recent publication has incorporated ancestry analysis using SNP genotyping to validate previously reported ethnicities of cell lines as well as elucidate the ancestral proportions of racially unidentified cell lines (23). These efforts are important and provide the scientific community with the (1) confirmation or rejection of the ethnicity designations assigned to these cell lines; (2) prevention of wasted time and money; (3) prevention of misleading publications leading to retractions; (3) confirmed ethnicity of previously unidentified cell lines; and (4) increased variation among cell lines.

In this study, we expand upon this recent focus and include 15 human cell lines representative of various cancer types for ancestry analysis. Using a set of 105 established Ancestry Informative Markers (AIMs) validated as SNP genotypes for population structure analyses, we assign respective West African (WA), Native American (NA), and European (EUR) ancestral proportions (36, 37). Key findings of our study include the high NA proportion found within the MDA-MB-468 breast cancer cell line suggesting this patient may have been an Afro-Hispanic/Latina female, although currently simply designated as Black. We also identified the 22Rv1 cell line, currently racially unidentified yet recently categorized as carrying mixed genetic ancestry in a study using 29 AIMs, as a predominantly EUR prostate cancer cell line (23). Furthermore, we conclusively establish that the E006AA-hT prostate cancer cell line, designated as African American, in fact carries majority EUR ancestry.

Materials and Methods

Cell culture

DUI145 (ATCC, Cat. # HTB-81), 22Rv1 (ATCC, Cat. # CRL-2505), and HeLa (ATCC, Cat. # CRM-CCL-2) cell lines were purchased from ATCC and grown in a humidified incubator with 5% CO₂ at 37°C in R.A. Kittles' laboratory. Cells were routinely tested for mycoplasma contamination using the PCR Mycoplasma Detection Kit (ABM, Cat. # G238). Cell lines were cultured in RPMI 1640 medium (ATCC, Cat. # 30-2001) supplemented with

10% FBS (ATCC, Cat. # 30-2020), penicillin–streptomycin (Gibco, Cat. # 15140122), and gentamicin (Fisher Scientific, Cat. # 15710064) as recommended by the supplier. Note that 0.2% Normocin (Invivogen, Cat. # ANT-NR-1) was added to the medium to prevent contamination by mycoplasma, bacteria, or fungi. In accordance with NIH guidelines concerning the authentication of key biological resources and in order to ensure the identity and validity of the resource, cell lines cultured in this study were purchased from ATCC that performs cell line characterizations. DNA was extracted from these cell lines within 2 weeks from receipt and not exceeding 3 passages. DNA extracted from the RWPE1 cell line was provided by L. Nonn. DNA extracted from the E006AA-hT cell line was tested from three separate sources, including an original purchase from ATCC, and were kindly provided by S. Ambs, A. Sreekumar, and S. Patierno. DNA extracted from the HCC1500, HCC1806, MCF-10A, MDA-MB-453, MDA-MB-468, and T-47D cell lines was provided by S. Kimbro. DNA extracted from the MDA-PCa-2b cell line was provided by S. Lloyd. DNA extracted from the RC-77T/E, LNCaP, and PC3 cell lines was provided by R. Mitra.

SNP genotyping

SNPs that were previously identified and validated for estimating continental ancestry information in admixed populations were selected to identify the AIMs (36–38). The AIMs panel consisted of 105 SNPs and were genotyped using the Sequenom MassARRAY genotyping platform with iPLEX chemistry according to the manufacturer's recommendations. iPLEX assays were designed utilizing the Sequenom Assay Design software, allowing for single-base extension designs used for multiplexing. DNA was isolated from the cell lines, and multiplex assays were performed to amplify 10 ng of genomic DNA by PCR. PCR reactions were treated with shrimp alkaline phosphatase enzyme to neutralize the unincorporated deoxyribonucleotide triphosphate. A post-PCR single-base extension reaction was performed for each multiplex reaction using concentrations of 0.625 μmol/L for low mass primers and 1.25 μmol/L for high mass primers. Reactions were diluted with 16 μL of H₂O, and fragments were purified with resin, spotted onto Sequenom SpectroCHIP microarrays (Agena Bioscience, Product 10500), and scanned by MALDI-TOF mass spectrometry. Individual SNP genotype calls were generated using Sequenom TYPER software, which automatically calls allele-specific peaks according to their expected masses. A genotype concordance rate of 99% was observed for all markers. Genotyping call rates that exceeded 98.5% were included in the analyses.

DNA ancestry analysis

Individual admixture estimates for each cell line were calculated using a model-based clustering method as implemented in the program STRUCTURE v2.3 (39). STRUCTURE 2.3 was run using parental population genotypes from WAs, EURs, and NAs (36) under the Admixture model using the Bayesian Markov chain Monte Carlo method and a burn-in length of 30,000 for 70,000 repetitions. Because we are unsure about the ancestries of our cell line samples, we used the admixture model to determine which estimation of K (number of sub populations) is the best fit for the data. We set K from 2 to 5 and ran 100 iterations. We determined that K = 3 had the best fit. We used the K = 3 estimates for our analyses.

Multidimensional scaling analysis

Multidimensional scaling (MDS) analysis was performed to visualize the genetic similarity of the cell lines to worldwide populations. Note that 1000 genomes variant data were downloaded, and 824 individuals representing 8 groups were selected to represent worldwide populations—Mende in Sierra Leone (MSL); Luhya in Webuye, Kenya (LWK); Toscani in Italia (TSI); British in England and Scotland (GBR); Indian Telugu in the UK (ITU); Han Chinese in Beijing, China (CHB); Chinese Dai in Xishuangbanna, China (CDX); and Gujarati Indian in Houston, TX (GIH; ref. 40). Markers matching the 105 AIMs were then selected for the MDS analysis. Individuals were removed with missingness > 0.05. Markers were removed with missing genotypes > 0.05 or minor allele frequency < 0.05. Though sparsely located across the genome, PLINK software was still used to remove markers in linkage disequilibrium using an r^2 greater than 0.5 in a 50 SNP window with a 5 SNP sliding window in the combined cell line and 1KG variant data (41). PLINK software was also used to perform the MDS analysis on the remaining 87 markers.

Results

One hundred five AIMs were selected from a larger previously validated set of markers to define critical genome candidate regions and characterize samples from diverse population groups (36, 37). This subset of AIMs contains specific SNPs capable of distinguishing WA, NA, and EUR genetic ancestry. Our ancestry estimates suggest cell lines previously classified as White/Caucasian by ATCC were accurately described, with mean EUR ancestry proportions of 97% (range, 92%–99%; Table 1). MCF-10A and MDA-MB-453 breast cancer cell lines were found to carry 97% and 99% EUR ancestry, respectively. PC3, DU145, LNCaP, and RWPE1 prostate cancer cell lines were found to carry 98%, 99%, 92%, and 96% EUR ancestry, respectively.

Similarly, results of some ancestry estimates for cell lines previously classified as Black/African American were accurately described. For example, the MDA-PCa-2b prostate cancer cell line was found to carry 86% WA ancestry (Table 1). Likewise, the HCC1500, HCC1806, and MDA-MB-468 breast cancer cell lines

were found to carry 79%, 80%, and 77% WA ancestry, respectively. Interestingly, the MDA-MB-468 breast cancer cell line was also found to carry an appreciable amount of NA ancestry (23%) suggesting possible Afro-Hispanic/Latina ancestry. Although the HeLa cervical cancer cell line was found to carry majority WA ancestry (66%), the WA proportion falls below the mean of approximately 80% WA ancestry typically observed in U.S.-born African Americans.

There are several cell lines that were included in our study that do not have racial identifiers specified by ATCC. For example, T-47D is a breast cancer cell line previously racially unidentified. Our ancestry analysis revealed that T-47D carries 100% EUR ancestry (Table 1). Similarly, the racial identification of the 22Rv1 prostate cancer cell line was never included within the biological characteristics specified when originally derived (42, 43). For this reason, no racial identifier has been available through ATCC (23, 24). Although a recent publication found this cell line to carry mixed genetic ancestry, our ancestry analysis revealed a majority EUR ancestry (99%; ref. 23).

Our genetic ancestry analysis of the E006AA-hT prostate cancer cell line revealed an extreme misclassification of racial identity compared with what has been reported in the literature as well as what is described by ATCC (24, 44). Although this cell line is sold commercially as an African American prostate cancer cell line, ATCC provides a disclaimer within the "Characteristics" section stating "during the accessioning of this line ATCC ran a short tandem repeat (STR) profile for the original starting material. The results match the characterization data in the cited references, however the STR profile was also found to match the STR profile (an 86% match) of another ATCC cell line, 786-O, a cell line derived from a renal cell carcinoma. The originating laboratory did not use the ATCC cell line, 786-O" (45). Due to the perplexity surrounding the true classification of this cell line, we included three separate DNA extraction samples provided by three separate laboratory groups of the E006AA-hT cell line. The SNP genotyping was found to be identical in all three E006AA-hT samples, revealing this cell line carries 91% EUR ancestry (Table 1).

We also included samples in our study of cell lines that are commonly used for research but are not commercially available.

Table 1. Ancestral proportions of commonly used human cancer cell lines

Cell line	ATCC catalog number	Gender	Age	Tissue	Designated race/ethnicity	WA	NA	EUR	Putative ancestry
PC3	CRL-1435	M	62	Prostate cancer	White/EA	0.3%	0.4%	99.3%	EA
DU145	HTB-81	M	69	Prostate cancer	White/EA	0.4%	0.5%	99.1%	EA
LNCaP	CRL-1740	M	50	Prostate cancer	White/EA	6.5%	1.9%	91.7%	EA
RWPE1	CRL-11609	M	54	Prostate cancer	White/EA	2.2%	1.7%	96.1%	EA
22Rv1	CRL-2505	M	N/A	Prostate cancer	N/A	0.3%	0.4%	99.3%	EA
E006AA-hT	CRL-3277	M	50	Prostate cancer ^a	Black/AA	1.3%	7.4%	91.3%	EA
MDA-PCa-2b	CRL-2422	M	63	Prostate cancer	Black/AA	86.5%	3.0%	10.6%	AA
RC-77T/E	N/A	M	63	Prostate cancer	Black/AA	88.9%	0.4%	10.7%	AA
HeLa	CCL-2	F	31	Cervical cancer	Black/AA	66.5%	0.4%	33.1%	AA
MCF-10A	CRL-10317	F	36	Breast disease	White/EA	0.9%	2.4%	96.7%	EA
MDA-MB-453	HBT-131	F	48	Breast cancer	White/EA	0.2%	0.4%	99.4%	EA
HCC1500	CRL-2329	F	32	Breast cancer	Black/AA	78.8%	0.5%	20.7%	AA
HCC1806	CRL-2335	F	60	Breast cancer	Black/AA	80.2%	4.4%	15.4%	AA
MDA-MB-468	HTB-132	F	51	Breast cancer	Black/AA	77.1%	22.7%	0.2%	AA/HA
T-47D	HTB-133	F	54	Breast cancer	N/A	0.1%	0.3%	99.7%	EA

Abbreviations: AA, African American; EA, European American; EUR, European; F, female; HA, Hispanic American; M, male; NA, Native American; N/A, not available; WA, West African.

^aE006AA-hT is sold by the ATCC as a prostate cancer cell line; however, ATCC acknowledges that the STR profile of this cell line is an 86% match to the 786-O renal cell carcinoma cell line (44).

For example, the RC-77T/E prostate cancer cell line was developed from an African American prostate cancer patient, and we confirmed the WA ancestral proportions of this cell lines at 89% (46).

As an additional method to verify the accuracy of cell line genetic ancestry characterization, MDS analysis was performed to visualize the genetic similarity of the cell lines against worldwide populations from the 1000 genomes project (Fig. 1). As expected, when plotting the cell lines and 1KG groups using the first two MDS dimensions, cell lines with predominantly EUR ancestry clustered with individuals from the GBR and TSI groups and admixed cell lines with predominantly WA ancestry clustered near the MSL and LWK groups. The first two MDS dimensions provide evidence that E006AA-hT is a cell line of predominantly European descent, due to its clustering with the European 1KG individuals, and the MDA-MB-468 cell line as possibly of Afro-Hispanic/Latina descent, due to its position on the axis between the WA and East Asian groups (a proxy for NA ancestry). As the MDA-MB-468 and HeLa cell lines are more heavily admixed, their genetic similarity lies between the East Asian to WA axis and between the EUR to WA axis, respectively.

Discussion

In order to promote meaningful and quality health disparities research, there has been a recent interest in incorporating increased racial diversity among human cell lines (17, 19, 23). Racial classification remains extremely useful for describing general patterns of health as most data are reported by self-identified race (47). However, we recognize that race also embodies social and cultural constructs, and most commonly used human cell lines were developed at a time when self-reported race was considered a sufficient demographic detail (47). As the role of

biological determinants in disease acquisition and progression becomes better defined, we cannot undercut the importance of individual genetic background (48–50). Because the use of cell lines is necessary to elucidate genes and pathways driving disease disparities, it is imperative to ascertain the accurate genetic ancestry of cell lines used in preclinical research in order to adequately explore the impact of genetic contributions on incidence and progression. Recognizing the importance of precise genetic assignment for research biospecimens, we sought to confirm or negate the current racial identification of commonly used cell lines, provide accurate and robust global ancestry estimates for cell lines from admixed individuals, while also revealing the genetic ancestry of previously racially unidentified cell lines.

Overall, our ancestry analysis mostly confirmed the racial classifications previously assigned to cell lines used in this study. In other words, most cell lines classified as "Caucasian/White" carried majority EUR genetic ancestry, whereas most cell lines classified as "African American/Black" carried majority WA genetic ancestry. However, a few key findings from our study refuted what has been reported in the literature and/or by ATCC. Most erroneously, we found E006AA-hT to carry 91% EUR genetic ancestry. This finding is problematic as E006AA-hT is currently marketed and commercially available as an "African American" prostate cancer cell line (24). Although ATCC provides a disclaimer that E006AA-hT matches the STR profiling of a renal cell carcinoma cell line, 786-O, there is no indication provided that the racial identifier of "African American" is not accurate (45). Because of the increasing mistrust of the E006AA-hT cell line we have observed in our interactions with others within the prostate cancer research community, we sought to include multiple samples of this cell line from three different laboratories located in distant geographical locations to firmly establish the true genetic ancestry. The implications of our finding are considerable as many laboratories have published or are in the process of publishing manuscripts exploring prostate cancer health disparities incorporating E006AA-hT as an African American prostate cancer cell line when, in fact, the cell line is neither African American nor likely derived from the prostate. The revelation of E006AA-hT as carrying predominant EUR ancestry is further disappointing as the field of prostate cancer health disparities research is already hindered by a lack of commercially available African American/Black cell lines, and this new EUR ancestral assignment leaves MDA-PCa-2b as the sole commercially available Black prostate cell line (23, 24).

Our study also highlights the advantage of utilizing an ample number of AIMs when conducting ancestry analysis. Recently, the 22Rv1 prostate cancer cell was found to carry mixed genetic ancestry using a smaller subset of 29 AIMs (23, 36). Although subsets of AIMs as small as 24 have been shown to be useful tools for ascertaining the origin of subjects from particular continents and correct for population stratification in admixed population sample sets, we suggest a more comprehensive subset of at least 100 AIMs as a large decrease in EUR performance has been observed in marker sets smaller than 64 (36). Using a validated set of 105 AIMs, we found 22Rv1 to carry 99% EUR ancestry. Because no racial identifier has ever been assigned to the 22Rv1 cell line, our genetic ancestry results clarify the ambiguity.

Given the high heterogeneity of African Americans, it is imperative to tease out the WA ancestral proportions of individual cell lines (51). We are the first to report that the commonly used HeLa cell line derived from African American cervical cancer patient Henrietta Lacks carries 66% WA ancestry (52). The average African

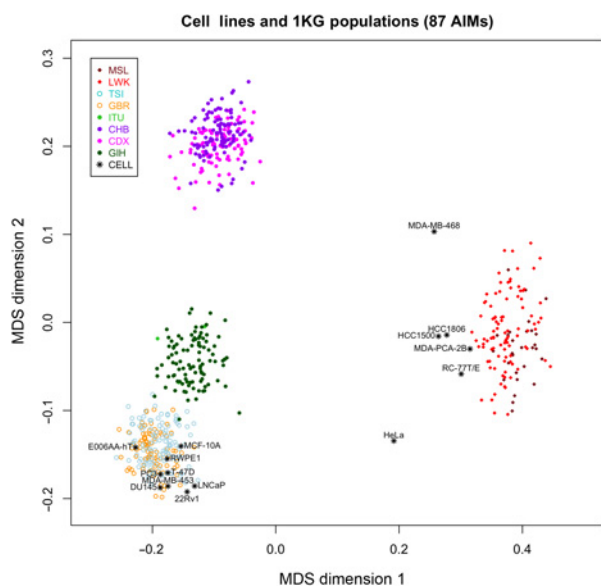


Figure 1. Genetic similarity of cell lines with worldwide populations. MDS analysis was used to determine the genetic similarity of the cell lines to groups in the 1000 genomes (1KG) project. Eighty-seven markers were used to generate MDS dimensions for 15 cell lines and 824 1KG individuals. Cell lines and 1KG groups are color-coded. Here, the East Asian group serves as a proxy for NA groups. CELL, a human cancer cell line.

American carries approximately 80% WA genetic ancestry, yet historically, African Americans have been likely to self-identify as Black regardless of how much or little European background they may possess (53). This self-identification as Black has followed the rule of hypodescent, under which any amount of Black ancestry warrants an association as African American (54, 55). Although the social and cultural influences of race on disease are undeniable, the role of genetics cannot be ignored. For example, an admixed individual with 20% EUR genetic ancestry may be at greater or lesser risk of certain diseases than an individual with 40% EUR genetic ancestry (56–60). In addition, the implications for pharmacogenomics exist as the nuances of drug effects or mechanisms may not be generalizable in the African American population due to high WA genetic variance (28, 61). For these reasons, there remains a burden on the scientific community to expand the collection of currently available human cell lines by incorporating more non-European options to capture the complete picture of genetic contribution to disease incidence, aggressiveness, progression, and response to treatment.

The need for increased diversity in research biospecimens is glaringly obvious when noting the near-complete lack of commercially available Hispanic/Latino cell lines (24). In our own ancestry analysis of the cell lines included in this study, we observed very low proportions of NA ancestry in all cell lines except the MDA-MB-468 breast cancer cell line. Although the MDA-MB-468 cell line is reported as Black, we found that it carries 77% WA ancestry and 23% NA ancestry (24). Based on these genetic ancestry proportions, it is reasonable to presume that the breast cancer patient from whom this cell line was derived may have been Afro-Hispanic/Latina as recent studies have highlighted this admixture proportion within a Hispanic-Caribbean population (62, 63). Thus, our genetic ancestry analysis may have uncovered an additional cell line with which to measure the impact of NA ancestry on breast cancer disease incidence, progression, and treatment.

As we progress further into an era of personalized medicine, the importance of racially diverse cell lines will grow clearer. Although it is imperative that research biospecimens are designated accurately in terms of race/ethnicity, it is crucial that they be characterized globally and locally for their genetic ancestry so that findings can be properly contextualized for the representative

populations. In the future, it would be ideal for commercial companies to report these global and local findings. In addition, genetic distance mapping of cell lines with the 1000 genomes or Human Genome Diversity Panel populations, using dimensional reduction techniques such as MDS or principal component analysis, should be performed to further determine the race/ethnicity of cell lines through their clustering with worldwide populations (40, 64). These techniques should further avoid misclassification that could occur with relying solely on AIMs designed to discern genetic ancestry proportions for a few discrete ancestral populations. We intend for the results of this study to encourage the scientific community to pursue ancestry analysis of additional cell lines as well as develop a wider range of diverse biospecimens.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Authors' Contributions

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