Establishment, authenticity, and characterization of cervical cancer cell lines

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ABSTRACT
Cell lines have been considered excellent research models in many areas of biomedicine and, specifically, in the study of carcinogenesis. However, they cease to be effective models if their behavior changes. Although studies on the cross-contamination of cell lines originating from different tissues have been performed, little is known about cell lines derived from cervical neoplasia. We know that high-risk HPV (HR-HPV) is associated with the development of this type of cancer. This link between HPV infection and cancer was first established over 35 years ago when HPV16 DNA was found to be present in a large proportion of cervical cancer biopsies. The present review paper aims to report the status of the establishment, authenticity, and characterization of cervical cancer (CC) cell lines. This is a systematic review of articles on the establishment, authenticity, and characterization of CC cell lines, published from 1960 to date in the databases and in cell repository databases. 52 cell lines were identified in the literature. Only 25 cell lines were derived from cervical neoplasia, of which only 45.8% have a reported identity test (genomic fingerprint). Despite the increase in the establishment of cell lines of cervical neoplasia and the standards for the regulation of these study models, the criteria for their characterization continue to be diverse.

Introduction

Even though cervical cancer (CC) may be preventable, it continues to occupy fourth place in the incidence of neoplasms worldwide, with 604,127 cases per year. In Mexico, it is the second most common cancer in the female population, with 9,439 cases per year.1 However, in middle and low-income countries, mortality can exceed 50% of cases. Strategies for prevention, timely diagnosis, and therapeutic interventions have been developed thanks to cancer research based on models using cell lines. The HPV life cycle is intrinsically linked to the programmed epithelial differentiation of the cervix. The majority of the focus has been on high-risk HPVs (HR-HPVs), which have been linked to cancer development.2,3 This link between HPV infection and cancer was first established over 35 years ago when HPV16 DNA was found to be present in a large proportion of cervical cancer biopsies.4–6 Cell lines are used in multiple areas of biomedicine, specifically in the study of carcinogenesis. A cell line is "an in vitro culture of cells that achieve indefinite survival and can be frozen and retrieved a theoretically infinite number of times."6 When the right conditions are met, these cells will keep dividing and keep many of the characteristics of the cell type or tissue from which they came. The European Collection of Animal Cell Cultures, 2012,8 defines two types of cell lines according to the capacity of the cells to continue dividing: finite or senescent, which die after a fixed number of duplications, and continuous, which can be propagated indefinitely. Every cell line comes from a specific tissue or by cloning an already established line; each tissue has specific characteristics (morphology, organization, specialized functions, cell cycle, etc.). When a cell line is established, many of these characteristics are lost, such as cell-cell (or cell-tissue) junction, cell communication, and regulation.7 The characteristics of a cell line vary from one person to another. However, they all share certain features. However, working with these models has a limited efficiency due to their behavior, i.e., this can change for two main reasons: contamination with other lines and misidentification.10 Multiple studies have demonstrated these issues, and it is estimated that between 18 and 36% of cell lines are contaminated or misidentified,11 with only 43% of cell lines considered "well identified.12 The cost of cell line contamination is high. For example, in the United States in 2013, investment in breast cancer research amounted to 370,000 million dollars, and it is estimated that at least 100 million dollars were lost in studies using cell lines that did not come from this tissue.13 In this case, not only are investment costs at stake, but also the credibility of the researchers and the inability to reproduce and transfer the results they come up with.

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Dr. Gartler published the first report of contamination in cell lines in 1968, demonstrated by electrophoretic analysis of the enzymes Glucose 6 Phosphate Dehydrogenase (G6PD) and Phosphoglucomutase (PGM). He evaluated 30 different cell lines, finding that 18 of them, despite coming from the Caucasian race, presented the same polymorphism (A) as the HeLa cells, which come from the African-American race, concluding that it was contamination of these lines and introducing the concept of biochemical polymorphisms. Since then, multiple studies corroborate his results have been conducted. Recently, the International Cell Line Authentication Committee (ICLAC) has reported 486 lines as misidentified, which has harmed at least 32,000 published articles.

Capes-Davis et al. (2012) point out that a cell line is misidentified when its DNA profile is no longer consistent with that of the donor from which it was first obtained. Other researchers corroborate this fact since there is an error when describing cell lines according to the tissue of origin. For example, the ECV304 line is said to come from normal endothelial cells when in fact it comes from bladder cancer. The KB line is said to come from laryngeal carcinoma when in fact it is a clone of modified HeLa cells that have been grown in a lab. It’s also possible to mix up culture samples, cross-contaminate cell lines with each other, and get biological contamination from microorganisms like mycoplasma and fungi.

Stacey (2007) indicates that there are three fundamental characteristics to ensure the quality of work with cell lines: 1) purity, that is, that they are free of microorganisms; 2) identity, which refers to the cells’ being who they say they are; and 3) stability, indicating that the genotype and phenotype must remain unchanged during growth and in vitro passages. Other criteria highlighted by cell culture researchers are the verification of viability, karyotyping, confirmation of the species of origin, specific cell identification (genomic fingerprinting), cell markers, genetic expressions, pluripotency (in the case of stem cell tissues), as well as quality controls in culture. Cell morphology ( phenotypic changes) and ploidy (genotypic changes) are also things to keep an eye on in cell lines to make sure they have the right biological properties.

The criteria for authentication and characterization of a line are very diverse among researchers. Therefore, international organizations such as the International Committee for Cell Line Authentication, the European Society for Animal Cell Technology, the World Biological Standards Institute, or the UK Cancer Research Coordinating Committee, among others, issue guidelines on the use, maintenance, development, and deposition of cell lines. Among the norms described by these organizations, the ethical aspects stand out, such as the need for the approval of the study by a research ethics committee, obtaining the informed consent of the donor, the transfer of rights to the biological material, etc. The data obtained on the tissue and the donor, the methods suggested to verify the authenticity of the cell line by DNA analysis, the criteria for its deposit in repositories, and even data on the establishment and characterization of the line should be included in the publications.

Also, the techniques used in the characterization and authentication of cell lines are varied. For example, isoenzyme analysis determines the origin of the species of a given line and allows the detection of cross-contamination through the electrophoretic mobility of different isoenzymes. Cytogenetic analysis allows chromosomal markers to be identified. Genomic fingerprinting exploits the variability found in non-coding regions of the human genome, which is organized into repeated sequences (variable numbers of VNTR tandem repeats) of two types: minisatellites (10 to 100 bp) and microsatellites (2 to 5 bp), also called short tandem repeats (STRs).

The analysis of these regions is necessary because the probability of two unrelated individuals having the same combination at a specific locus is less than 1%. In addition, advances in technology have favored the evaluation of cell lines. From morphological observation of cells in culture by transmission or scanning electron microscopy, to the implementation of genomic analysis methods and molecular cytogenesis, researchers now have more accurate methods to assess the authenticity of cell lines and their growth characteristics in culture.

In summary, studies on cross-contamination of cell lines originating from diverse tissues have been performed, and the development of these biological models has increased significantly in recent decades. However, despite this, there is a lack of information on appropriately identified cervical neoplasia-derived cell lines. So, this review will show how cervical cancer (CC) cell lines are made, how they are authentic, and how they are characterized.

**Target**
To show how cervical cancer (CC) cell lines have been set up, authenticated, and characterized over time.

**Materials and methods**
The present study summarizes the current knowledge concerning the establishment and characterization of CC cell lines established between 1960 and 2020. It is a systematic review based on the phases of the Preferred Reporting Items for Systematic Reviews and Partial Analyses (PRISMA). The concepts and strategies for the systematic search to locate information from Pozos and Garrocho (2012) were also taken up.

A search for publications concerning the establishment and characterization of CC cell lines from 1960 to 2020 was performed in the databases PubMed, Web of Science, Natura, SpringerLink, EBSCO, ScienceDirect, Ovid, Redalyc, PLOS, BMC, SciELO, PMC, Google, Google Scholar, BIG (Search for Global Information) and Academic Source. These cell repositories, including the American Type Culture Collection (ATCC), European Collection of Authenticated Cell Cultures (ECACC), Accegen, and Celsolus, were also looked at.

All titles that presented at least one of the following keywords: establishment, characterization, authenticity and cervical cancer cell lines were reviewed. Initially, papers that did not respond to any of the keywords, articles written in languages other than Spanish, English, or French, and titles without an abstract or text were eliminated. Next, abstracts of articles on
the establishment and/or characterization of CC cell lines were obtained, excluding those in which the lines had originated from animal models, from other already established lines (i.e., by cloning, transfection, or genotypic modification), or those derived from Low-Grade Intraepithelial Lesions (LGEI).

Subsequently, references to CC lines were obtained and analyzed for their integration or exclusion in the study. The HeLa cell line was first ruled out because it had been reported as a contaminant of other cell lines and because no specific papers on how it was made or how it was characterized were found. However, many papers on some of its characteristics were found.

The documents were then obtained in PDF format, and the data was organized under two headings: 1) the establishment of CC cell lines and 2) the characterization of the lines. An EXCEL database was constructed considering the following: year of publication, authors, journal, names of the established cell lines, general data (age, race, histopathological diagnosis, tissue origin of the sample, previous treatment), characterization methods (morphological, growth characteristics, cell population, doubling time, contact inhibition, adhesion, saturation density, karyotype, HPV genotyping, isoenzymatic analysis, genomic fingerprinting), and ethical aspects.

To verify that the lines were derived from cervical neoplasia, they were examined for: a) karyotype, b) HPV genotyping (due to its association with cervical carcinogenesis), and c) tissue of origin. A second document review of each line was performed to identify missing data regarding these three conditions. The data on the methods used to verify the authenticity and characterization of each line were subsequently organized into tables (Figure 1). Finally, the results, according to their classification, were expressed in percentages.

**Results**

A total of 378 abstracts were reviewed from which 140 scientific articles were obtained. In these papers, the establishment of 52 CC cell lines from 1960 to 2020 was reported.

In terms of origin, we found that the countries reporting the highest percentage (53.7%) of established lines are in Asia, followed by North America (24.1%) and Latin America (11.1%), while European countries report the lowest percentage (9.3%) of established lines. Only one of the lines (Do'Tc2) did not report this data. Table 1 shows data that was consistent across all reviewed papers on the establishment and characterization of CC cell lines in the study period. Only 48% of the 52 cell lines were identified as derived from cervical neoplasia as they come directly from cervical tissue and have karyotyping and HPV genotyping reported as methods to assess, respectively, authenticity and determine the main characteristics of the line. The remaining lines (52%) lack proper identification because: (a) they are derived from metastatic tissues (lymph nodes-HT3, MS751-), intestinal epithelium-CaSki-, omentum-ME180-) or cells derived from fluids such as ascitic fluid (EC82, SFCC, SKS), b) no karyotype report (EC82), no genotyping (SFCC), or c) no general data report (age).

According to Table 2, which contains the data of the 25 CC lines identified according to their racial origin, the following results were obtained: 10 of them (40%) come from Caucasian women, 13 (52%) from Asian women, and only 2 (8%) from Latin American women. Many lines (40%) correspond to women between 30 and 39 years of age, followed by those obtained from women between 40 and 49 years of age (28%) and those between 60 and 69 years of age (20%). A few lines (8%) derive from women in their 50s and just one line does not report the data (INBL).

Regarding the type of neoplasm, 76% of the lines are described as epithelial carcinomas and only 12% as adenocarcinomas (×H1, SiSo, RSBS-43). 8% of the authors do not specify whether it is squamous or glandular (CA and SKG-III), and only one line (4%) is said to come from a lymphoepithelial type cancer (HUUCLE) determined by histology. Concerning the degree of progression of the neoplasm, we found great variability among the authors since the majority (80%) only refer to the stage according to the FIGO classification and only 4% include the grade, while 16% do not refer to the progression of cancer. The lines classified as stage I occupied 28% of the total, those included in stages II and III were 20%, and only 16% came from metastatic cervical neoplasms (stage IV).

Because of the association of cervical cancer with high-risk HPVs, viral genotyping has become an important element in cell line specificity. Of particular interest for study in carcinogenesis are those that have been reported: a) positive for viral load and b) infected with more than one HPV type most of the lines (56%) are described as HPV16, followed by those characterized as HPV18 (C4, SW756, CaLo, INBL, SiSo) and HPV negative (C33, CX, CA). Only one line was reported as HPV59 (HHUS) and one as HPV56 (HUUCLEC). In the line named CUMC-6, both type 16 and type 18 viral genomes were identified.

All lines are tumorigenic, so the karyotypes showed aberrations in different chromosomes and were referred to as aneuploid (60%), triploid (12%), diploid or hyperdiploid (8%), respectively. A few have been described with other chromosomal alterations, such as SiHa (hypermultiploid), CUMC-3 (hypotetraploid), and, most notably, CX, which was reported as euploid (46 chromosomes), despite structural alterations in the XX chromosome.

Table 3 indicates the data reported for the characterization of each line; these include those concerning 1) culture conditions (cell division time, growth density, contact inhibition), 2) confirmation of line identity (genomic fingerprinting, isoenzyme analysis, confirmation of tissue of origin) and 3) confirmation of non-contamination.

**Table 1.** Cervical cell lines established between 1960–2020.

| Criteria | No. cell line identified | Percentage |
|----------|--------------------------|------------|
| Not reported karyotype | 13 | 25.0 |
| Not reported genotyping HPV | 3 | 5.8 |
| Ciudad de México | 7 | 13.5 |
| Not reported general date or karyotype/ genotyping | 4 | 7.7 |
| Karyotyping + Genotyping HPV | 25 | 48.0 |
| Total | 52 | 100 |
GENERAL OUTLINE OF THE STUDY

Articles identified in search engines
n=78,464

Articles identified with the inclusion criteria
n=378

Excluded articles: not corresponding to the topic, cell lines originating from other tissues, foreign languages (German, Japanese, Chinese), without abstract or full text. N=78,086

Articles excluded they correspond to derived cell lines: from animal models, from already established lines (cloned, transfected or modified), from low-grade lesions. n=238

Summary of articles for your review
n=140

Identification of Cervical Cancer cell lines established in the study period
n=52

Cervical Cancer cell lines characterized only by HPV karyotyping and genotyping
n=25

Excluded cell lines: not having general data, coming from metastatic tissues, without HPV karyotype and / or genotyping. No informed consent, lack of proper documentation. n=63

Figure 1. General outline of the study.

Although many of the data reported in the studies were in general, some of the most constant ones were the time in which the cell population in culture doubles its population, as well as the confirmation of the tissue of origin. However, a considerable percentage (24%) of the total lines report 5 or fewer characteristics (some only two-TC140-); the majority (68%) include 6 to 7 criteria in their characterization (including authenticity and free of microorganisms), while only 8% include 8 to 9 (SKG-III and SiSo, respectively).

Among the lines with accelerated growth (24 h) were XH1, CX, and CA, followed by those with intermediate growth (24–48 h), which corresponded to 48%, while 24% of the lines presented slow growth times (48–76 h). Two lines (8%) - CaLo, INBL-were reported with the slowest cell division times (more than 76 h) and two others (TC140, TC146) did not report the data. The number of times cells were subcultured varied from line to line, from 34 passages (SiSo) to 300 (CUMC-6); only a few lines (SiHa, CX, TC140, TC146, RSBS9, RSBS14, RSBS23, RSBS43) do not have the report.

One of the criteria rarely found in the publications was the inhibition of cell growth by contact. Only in 11 lines (44%) was this feature described; in seven of them their cells do not stop dividing upon contact with others, as happens in neoplastic tissues, while in four lines (RSBS-9, RSBS-14, RSBS-23, and RSBS-43) contact inhibition was documented. Regarding the density at which the cells grow, in 36% (9 cell lines), this characteristic has been reported using qualitative (high, low) and quantitative (various) scales as shown in the table.

On the other hand, the characterization of tumorigenic capacity was established in 21 lines (84%); the privileged model for this was murine (18 lines), followed by 3D cultures (4 lines), while in three cell lines (CaLo, INBL, and CX) no publications were found; only one line (HUUCLEC) reported the absence of carcinogenesis in a murine model after inoculation. Regarding the establishment of line specificity and identity, methods based on DNA analysis (SRT, VNTR, SNP, FISH) were recurrent (52%) among the investigators (13 lines), whether used alone or in combination with RNA analysis (transcriptome) or protein expression (proteome). However, in only 44% of the lines, no other genetic tests were done other than karyotyping.

Likewise, the isoenzymatic analysis of biochemical polymorphisms seems to be a common criterion for the lines established before the 1990s, since in 11 lines (44%), established in that period, this data was documented, while for the rest it was not; the most studied enzymatic polymorphisms were in G6PD, PGM1 and 3, AK-1 and ACP1. The only line that didn’t report on this was C33. All the other lines reported on either a) specific epithelial tissue antigens (vimentin, desmogevin, epithelial membrane antigen, etc.), specific neoplastic tissue antigens (carcinoembryonic antigen), histochemical or morphological studies, or a combination of these.
| No. | Cell line | Histology diagnosis | Race | Age | HPV | Karyotype | Reference |
|-----|-----------|---------------------|------|-----|-----|-----------|-----------|
| 1   | C4        | Exophytic invasive squamous carcinoma of the cervix stage II, grade IV | C    | 41  | HPV18 | Hyperdiploidy | Auersperg, Friedl |
| 2   | C33       | Invasive cervical carcinoma grade IV | C    | 66  | Negative for HPV | Hyperdiploidy | Auersperg, Friedl |
| 3   | SiHa      | Squamous cell carcinoma stage II | A    | 55  | HPV16 | Hypertetraploidy | Kelland; Spencer; Han; Friedl |
| 4   | SW756     | Poorly differentiated invasive squamous carcinoma of the uterine cervix | C    | 46  | HPV18 | Triploid | Friedman; Popescu; Nozawa; Shirasawa; Kelland; Spencer; Han; Monroy |
| 5   | SKGII     | Uterine cervical cancer stage II | A    | 38  | HPV16 | Aneuploidy | Kelland; Spencer; Han; Monroy |
| 6   | HX151c    | Poorly differentiated squamous cell carcinoma stage IB | C    | 30  | HPV16 | Aneuploidy | Kelland; Spencer; Han; Monroy |
| 7   | HX155c    | Moderately differentiated squamous cell carcinoma stage IB | C    | 44  | HPV16 | Aneuploidy | Kelland; Spencer; Han; Monroy |
| 8   | HX156c    | Poorly differentiated squamous cell carcinoma stage IB | C    | 31  | HPV16 | Aneuploidy | Kelland; Spencer; Han; Monroy |
| 9   | HX160c    | Moderately well-differentiated squamous cell carcinoma stage IB | C    | 44  | HPV16 | Aneuploidy | Kelland; Spencer; Han; Monroy |
| 10  | XH1       | Invasive focally keratinizing adenosquamous | C    | 32  | HPV16 | Aneuploidy | Kelland; Spencer; Han; Monroy |
| 11  | CaLo      | Epidermoid cervical carcinoma of keratinized large cell from nonmetastatic tumor stage IIIB | L    | 55  | HPV18 | Aneuploidy | Kelland; Spencer; Han; Monroy |
| 12  | INBL      | Epidermoid cervical carcinoma of keratinized large cell from metastatic tumor stage IVA | L    | NR  | HPV18 | Aneuploidy | Kelland; Spencer; Han; Monroy |
| 13  | CUMC-3    | Invasive nonkeratinizing squamous cell carcinoma stage IIB | A    | 32  | HPV16 | Hypotetraploidy | Mark |
| 14  | CUMC-6    | Squamous cell carcinoma of the cervix uteri | A    | 31  | HPV16-18 | Diploid/trisomy | Kim |
| 15  | TC-140    | Moderately differentiated squamous cell carcinoma with metastases to lymph nodes | C    | 39  | HPV16 | Triplet (61) | Braun; Mark |
| 16  | TC-146    | Moderately differentiated epidermoid carcinoma in situ of the cervix | C    | 36  | HPV16 | Triplet (48) | Mark |
| 17  | SiSo      | Adenocarcinoma of the uterine cervix stage IB | A    | 67  | HPV18 | Aneuploidy | Sonoda; Chou |
| 18  | CK        | Squamous cell carcinoma of the uterine cervix stage IIA | A    | 48  | Negative for HPV | Euploidy | Kim |
| 19  | CA        | Non-keratinizing squamous cell carcinoma IB2 | A    | 36  | Negative for HPV | Aneuploidy | Isaka |
| 20  | HHUS      | Uterine cervical keratinizing squamous cell carcinoma | A    | 64  | HPV59 | Diploid (48) | Ishiwatari |
| 21  | RSB-9     | Moderately differentiated keratinizing squamous cell carcinoma stage III | A    | 49  | HPV16 | Aneuploidy | Javed |
| 22  | RSB-14    | Moderately differentiated non-keratinizing squamous cell carcinoma stage III | A    | 34  | HPV16 | Aneuploidy | Javed |
| 23  | RSB-23    | Poorly differentiated non-keratinizing squamous cell carcinoma stage III | A    | 45  | HPV16 | Aneuploidy | Javed |
| 24  | RSB-43    | Moderately differentiated adenosquamous of uterine cervix stage III | A    | 63  | HPV16 | Aneuploidy | Javed |
| 25  | HUUCLEC   | Human uterine cervical lymphoepithelial carcinoma | A    | 61  | HPV56 | Aneuploidy | Kirguchi |

C= Caucasian, A= Asian, L= Latin American, NR= Not reported data.

On the contrary, and even though the problem of cross-contamination between cell lines, as well as contamination by biological organisms, has been exposed since 1968, in nine lines (36%) no evidence was found for the absence of intra-or interspecies contamination, in 12 lines (48%) only contamination with mycoplasma or with another microorganism (Epstein-Barr virus) was evaluated, in three (12%) the absence of contamination with HeLa cells was reported, and only in one line (SW756) contamination with mycoplasma and HeLa cells were evaluated together.

Finally, in 76% of the lines identified, no ethical aspects were mentioned although the inclusion of these aspects has become an indispensable requirement in the publication of studies on the establishment of cell lines, in the procurement and use of tissues, as well as in the manipulation of genetic material. Likewise, ethical criteria are particularly important concerning the procurement of biological tissues, the transfer of rights, and the approval of the study by an ethics committee.

**Discussion**

**Establishment of cervical cancer cell lines**

78.8% of cervical cancer cell lines reported between 1960 and 2020 emerged from developed countries, such as Canada, the USA, Japan, the United Kingdom, and South Korea. A likely consequence of its high investment in research and development (R & D), however, is that the USA is the nation that invests the most in this concept (28.1% of national gross expenditure), followed by the European Union (19%) and Japan (10%). The remaining lines (21.2%) were developed in countries considered in transition (Mexico, Brazil, Thailand), where R & D investment is not so strong. 61

However, only 48% of the lines that were reported came from cervical tissue, so they can be used as biological models of how malignant cells behave in real life. Culturist researchers say that it is important to know where the cell line came from so that future research can show how the system model works. 62,63
Table 3. Methods to evaluate the authenticity and characterization of cervical cell lines.

| No. | Cell line | T_D | No. Passages | Contact inhibition | Saturation density | Tumorigenicity | Uniqueness | Isoenzyme analysis | Contamination | Confirmation the tissue of origin |
|-----|-----------|-----|--------------|--------------------|-------------------|----------------|------------|-------------------|--------------|----------------------------------|
| 1   | C4        | 24-48h | 90           | No                 | NR                | Collagen gel, hamster model | SRT, transcriptome | G6PD-B          | Free of HeLa       | Cytology, histochemistry,          |
| 2   | C33       | 24-76h | 70           | No                 | NR                | Murine model       | SRT, transcriptome | G6PD-B, PGM1–1, AK-1 | Free of HeLa       | NR                                |
| 3   | SiHa      | 2.6 days | 100         | NR                | NR                | Murine model       | SRT, transcriptome | G6PD-B, PGM1–1, AK-1 | Free of mycoplasma, or HeLa | Desmosomes, tonofilaments       |
| 4   | SKG-III   | 1.6 days | 80          | 4x10^9 cell/cm^2 | Murine model      | SRT, transcriptome | SRT, transcriptome | G6PD-B, GLO-1,2 | Free of mycoplasma, HeLa, ME180, or C4I | Free of HeLa | Desmosomes, tonofilaments, HLA-A 2402 |
| 5   | HX15c     | 42h    | 70           | NR                | Low               | Murine model       | G6PD-B, PMG1–1, PMG3-a, AK-1, PMG3-b | Free of mycoplasma, or HeLa | Vimentin, cytokeratin |                                    |
| 7   | HX15c     | 48h    | 50           | NR                | Low               | Murine model       | G6PD-B, PMG1–1, PMG3-a, AK-1, PMG3-b | Free of mycoplasma, or HeLa | Vimentin, cytokeratin |                                    |
| 8   | HX15c     | 30h    | 75           | NR                | High              | Murine model       | G6PD-B, PMG1–1, PMG3-a, AK-1, PMG3-b | Free of mycoplasma, or HeLa | Cytokeratin |                                    |
| 9   | HX16oc    | 68h    | 50           | NR                | Low               | Murine model       | G6PD-B, PMG1–1, PMG3-a, AK-1, PMG3-b | Free of mycoplasma, or HeLa | Vimentin, cytokeratin |                                    |
| 10  | XH1       | 16.2h  | 100          | NR                | NR                | Murine model       | SRT, transcriptome | MS1              | Free of mycoplasma, HeLa, A431, Calk, and Bowes melanoma | NR | Antibodies to membrane antigen, carcinoembryonic antigen, desmin, vimentin and cytokeratin Desmogelein-1 |
| 11  | CaLo      | 3–4 days | 50          | High (>15,000 cell/ml) | NR                | NR                | NR                | NR                | NR | NR | Desmogelein-1, HLA-A11 |
| 12  | INBL      | 3–4 days | 50          | High (>15,000 cell/ml) | NR                | NR                | NR                | NR                | NR | NR | Desmogelein-1, HLA-A11 |
| 13  | CUMC-3    | 48h    | 310          | No                 | NR                | Murine model       | SRT, transcriptome | G6PD-B, LDH-4, PGM | Free of HeLa       | Desmosomes, tonofilaments, HLA-DRB1 0401 |
| 14  | CUMC-6    | 36h    | 300          | No                 | NR                | Murine model       | SRT, transcriptome | G6PD-B, LDH, PGM | Free of HeLa       | Desmosomes, tonofilaments, HLA-DQw3 |
| 15  | TC-140    | NR     | NR           | 1x10^5 cell/cm^2   | Murine model with lung metastases | Murine model | NR | NR | Free of mycoplasma |
| 16  | TC-146    | NR     | NR           | 1x10^5 cell/cm^2   | Murine model       | SRT, transcriptome | G6PD-B, LDH, PGM | Free of mycoplasma |
| 17  | SiSo      | 24-35h | 34           | NR                | Not tumorigenic    | RPPA, SNP, transcriptome | NR | NR | Free of mycoplasma |
| 18  | CX        | 20h    | NR           | NR                 | Not tumorigenic    | RPPA, SNP, transcriptome | SRT | NR | NR | Free of mycoplasma |
| 19  | CA        | 14.3h  | 280          | NR                 | NR                 | Murine model       | NR | NR | NR | Free of virus Epstein y Barr |
| 20  | HUHUS     | 67h    | 70           | 3.9x10^4 cell/cm^2 | Murine model | NR | NR | NR | NR | Desmosomes, tonofilaments, carcinoembryonic antigen, cytokeratin |
| 21  | RBS5-9    | 48h    | NR           | Yes                | NR                 | Tumorosphere       | VNTR | NR | Free of mycoplasma |
| 22  | RBS5-14   | 48h    | NR           | Yes                | NR                 | Tumorosphere       | VNTR | NR | Free of mycoplasma |
| 23  | RBS5-23   | 48h    | NR           | Yes                | NR                 | Tumorosphere       | VNTR | NR | Free of mycoplasma |
| 24  | RBS5-43   | 48h    | NR           | Yes                | NR                 | Tumorosphere       | VNTR | NR | Free of mycoplasma |
| 25  | HU大切中 | 70h    | 60           | No                 | Not tumorigenic    | NR | NR | NR | Free of virus Epstein y Barr |

1. Doubling time, NR: Not reported data, STR: Short Tandem Repeats, SNP: single-nucleotide polymorphisms, VNTR: variable number of tandem repeats, M531: Hypervariable minisatellite probing, FISH: fluorescence in situ hybridization, RPPA: reverse phase protein arrays, ACP-1: human red cell acid phosphatase, AK-1: adenylate kinase polymorphism, G6PD: glucose 6 phosphate dehydrogenase, PGM: phosphoglucomutase, HLA: human leukocyte antigen
Different cervical cancer cell lines are important because they allow us to know how the cells will react to a stimulus or treatment when they are grown in xenograft cultures in vivo. This is because we need to know how the cells will react to a stimulus or treatment when they are grown in xenograft cultures.

In another respect, 9 of 25 cervical cancer cell lines were established between 1960 and 1990, so the obtaining of the cells was understood as an extension of the diagnostic process in which the patient did not need to be consulted. According to international organizations, only one cell line had its origin confirmed by the donor. At least 12 cell lines, established after the 1990s, lack an ethical review, i.e., obtaining biological material and consent or rights from the donor. In this case, if an ethics committee had given its approval to the methods for getting the tissue, that would be something that is not talked about.

**Authentication of cervical cancer cell lines**

The American National Standards Institute (ANSI) and the American Type Culture Collection (ATCC) and other international agencies (UKCCR, ECACC) promote the use of genomic analysis techniques (STRs, VNTS, minisatellites, etc.) for identity verification of human cell lines (ASN-0002) that are unique and unrepeatable. STR analysis is the most common method used by the authors. It is a standard method that is fast, cheap, and has been tested by cell banks. The data is also consistent.

Other techniques that corroborate the uniqueness of cell lines are HPV integration sites, microsatellite stability, HLA typing, polymorphisms, as well as mutations and gene expression. In our study, few researchers reported these latest data. One disadvantage of this is that some cell lines tend to undergo genetic changes with continuous passages in culture. In malignant cells, there may be loss of heterozygosity and increased instability of microsatellites.

Also, persistent high-risk papillomavirus infection, i.e., HPV16, has been significantly associated with neoplastic progression. However, the determinants of viral persistence and clearance are not yet well understood. Many studies have been done to compare the responses of different types of CC cell lines to the administration of antineoplastic substances, but the most common models have been genotyped with HPV16 or 18. This means that little is known about how neoplastic cells react to another or no viral type.

In recent studies conducted in Europe, North America, and Latin America, in more than 1000 women with positive samples for HPV, they show the presence of multiple infections at high and low risk in more than 30% of the positive cases studied. Multiple infections appear to be more common in women under the age of 30, whereas viral type 16 excels in single and multiple infections associated with high-and low-risk HPV. A single viral genotype has been detected in a single sample. Some studies have shown that having a lot of HPV can make it more likely that you'll get high-grade lesions and get cancer, but they aren’t sure yet.

In our review, a single cell line reports the viral presence of both HPV16 and 18 in a woman over 30 years of Asian origin. The development of models with these characteristics could help us learn more about how many infections affect cervical carcinogenesis, which would help us figure out who is at risk in groups.

At least five lines of cervical cancer do not report the data and, although few studies on cell line contamination include these five lines of cervical cancer, their data are not conclusive if these lines were found or not contaminated; the International Committee for Authentication of Cell Lines, reports five lines of CC that have been found as contaminants of other-HeLa, ME180, SKGII, C33A, and TCO2-.. The above data shows that it’s important to report how to keep cultures healthy and cell lines true.

**Characterization of cervical cancer cell lines**

Knowing the morphological characteristics of cells in culture allows identifying changes in response to different environmental conditions (i.e., changes in the substrate, cryopreservation, cell density, phenotypic and genotypic changes, etc.) that require constant monitoring to prevent the invalidation of research work.

Normal cells usually stop dividing at a high cell density. They block in the G1 phase of the cell cycle and deteriorate very little. Tumor cells result in morphological deterioration but continue to proliferate beyond confluence. Freshner suggests expressing cell density in cell/cm2 to avoid ambiguities in interpretations such as high, medium, and low of the correct seeding density and subculture interval is done by performing a growth curve to establish the deviations from this pattern and prevent cell deterioration.

Likewise, chromosome content in cancer lines is aneuploid (abnormal chromosome content) and heteroploid (variability in chromosome number in the same cell population) as a result of alteration of tumor suppressor proteins, i.e., p53 and Rb, leading to genetic instability throughout the subcultures. Viability and karyotype have also been used to check the genetic stability of cell cultures and to see if cells have changed.

Phenotypic changes can occur during different passages of the cultures or by intra-and interspecies contamination, so it is important to report the number of passages that have had the established line, as well as the purity tests used in the laboratory. Karyotype is the most commonly used method to monitor the genetic instability of culture and determine the malignancy of the cell line, yet almost half of the identified lines do not report it.

Ultimately, the results of the study indicate a significant increase in the establishment of cervical cancer cell lines in recent decades. However, despite the regulations of international agencies, authentication and characterization criteria remain very diverse among researchers and some dates, i.e., ethical review, are even unaffordable for the majority.

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Data availability statement

The authors hereby confirm that the data supporting the conclusions of this study are available in supplementary material.

References

1. International Agency for Research on Cancer. Cervix uteri. Source: Globocan 2020. World Health Organization. Disponible en 2021. In: https://gco.iarc.fr/today/data/factsheets/populations/900-world-fact-sheets.pdf
2. Doorbar J, Egawa N, Griffith H, Kranjec C, Murakami I. Human papillomavirus molecular biology and disease association. Rev Med Virol. 2015;25(Suppl 1):2–23. doi:10.1002/rmv.1822.
3. Cubie HA. Diseases associated with human papillomavirus infection. Virology. 2013;445(1–2):21–34. doi:10.1016/j.virol.2013.06.007.
4. Durst M, Gissmann L, Ikenberg H, zur Hausen H. A papillomavirus DNA from a cervical carcinoma and its prevalence in cancer biopsy samples from different geographic regions. Proc Natl Acad Sci U S a. 1983;80(12):3812–3815. doi:10.1073/pnas.80.12.3812.
5. Ja S, Patterson MR, Morgan EL, Macdonald A. The human papillomavirus oncoproteins: a review of the host pathways targeted on the road to transformation. J Gen Virol. 2021;102(3):001540. doi:10.1099/jgv.0.001540.
6. Alcántara LQ. Control de calidad en los cultivos celulares para ingeniería de tejidos (diapositivas en pdf). México: Universidad Autónoma de San Luis Potosí; 2017.
7. Freshney RI. Culture of animal cell: a manual of basic technique and specializes applications. 6th ed. USA: Wilby-Blackwell; 2010.
8. Colección Europea de Cultivos de Células Animales. 2012. European Collection of Authenticated Cell Cultures (ECACC). Fundamental techniques in cell culture. Laboratory handbook, 2th ed. 2
9. Langdon SP. Characterization and authentication of cancer cell lines: an overview. Methods Mol Biol. 2004;88:33–42. doi:10.1385/1-59259-406-9:9:3.
10. American type Culture Collection. ATCC® animal cell culture guide: tips and techniques for continuous cell lines. 2014. Disponible en: http://www.uaab.cat/doc/ATCCguide. Acceso el 31 de mayo de 2018.
11. Hughes P, Marshall D, Reid Y, Parkes H, Gelber C. The costs of using unauthenticated, over-passaged cell lines: how much more data do we need? BioTechniques. 2007;43(S):575–586. doi:10.2144/000112598.
12. Vasilevsky NA, Brush MH, Paddock H, Pointing L, Tripathy SJ, LaRocca GM, Haendel MA. On the reproducibility of unique identification of research resources in the biomedical literature. PeerJ. 2013;1:e148. doi:10.7717/peerj.148.
13. Freedman LP, Gibson MC, Ethier SP, Soule HR, Neve RM, Reid YA. Reproducibility: changing the policies and culture of cell line authentication. Nat Methods. 2015;12(6):493–497. doi:10.1038/nmeth.3403.
14. Gartler SM. Apparent HeLa contamination of human heterologous cell lines. Nature. 1968;217(5130):750–751. doi:10.1038/217750a0.
15. Nelson-Rees WA, Flandermeyer RR. Inter- and intraspecies contamination of human breast tumor cell lines HBC and BrCa5 and other cell cultures. Science. 1977;195:1343–1344. doi:10.1126/science.557237.
16. Nelson-Rees W, Daniels D, Flandermeyer R. Cross-Contamination of cells in culture. Science; 1981; 212(4493):446–452. doi:10.1126/science.6451928.
17. Drexel HG, Dirks WG, MacLeod RAF. False human hematopoietic cell lines: cross-contamination and misinterpretations. Leukemia. 1999;13(10):1601–1607. doi:10.1038/sj.leu.2401510.
18. Stacey GN, Masters JRW, Hay RJ, Drexler HG, Macleod R, Freshney RJ. Cell contamination leads to inaccurate data: we must take action now. Nature. 2000;403(6768):356. doi:10.1038/3500394.
19. Macleod R, Dirks WG, Drexler HG. Persistent use of misidentified cell lines and its prevention. Genes Chromosomes Cancer. 2002;33(1):103–105. doi:10.1002/gcc.1217.
20. Korch C, Spellman MA, Jackson TA, Jacobsen BM, Murphy SK, Lessey BA, Jordan VC, Bradford AP. DNA profiling analysis of endometrial and ovarian cell lines reveals misidentification, redundancy and contamination. Gynecol Oncol. 2012;127(1):241–248. doi:10.1016/j.ygyno.2012.06.017.
21. Peavey J, Malek G. Cell line authentication in vision research and beyond: a tale retold. Ophthalmo Vis Sci. 2020;61(6):19–25. doi:10.1167/iovs.20-26169.
22. Capes-Davis A, Reid Y, Kline MC, Storts DR, Strauss E, Dirks WG, Drexler HG, MacLeod RAF, Sykes G, Kohara A, et al. Match criteria for human cell line authentication: where do we draw the line? Int J Cancer. 2012;132:2510–2519. doi:10.1002/ijc.27931.
23. Dirks WG, MacLeod RA, Drexler HG. ECV304 (endothelial) is really T24 (bladder carcinoma): Cell line cross contamination at source. Vitro Cell Dev Biol. 1999;35(10):558–559. doi:10.1073/s11626-999-0091-8.
24. Buhring GC, Eby EA, Eby MJ. Cell line cross-contamination: how aware are mammalian cell cultures of the problem and how to monitor it? Vitro Cell Dev Biol Animal. 2004;40(7):211–215. doi:10.1209/1437-076X/40/4-2004/111.CHLHAA.2.CO.2.
25. Schweppre R. Thyroid cancer cell line misidentification; an update. J Clin Endocrinol Metab. 2013;98(3):956–957. doi:10.1210/jc.2012-4182.
26. Buhring GC, Valeco M, Pan CY. Cell culture contamination by mycobacteria. Vitro Cell Dev Biol Anim. 1995;31(10):735–737. doi:10.1007/BF02634111.
27. Stacey GN, Auerbach JM. Quality control procedures from stem cell lines. Culture of human stem cells. Freshney R, Stacey G and Auerbach J, editors. John Wiley & Sons, Inc; 2007.
28. Nims RW, Reid Y. Best practices for authenticating cell lines. Citro Cell Dev Biol Animal. 2017;53(10):880–887. doi:10.1126/s11626-017-0212-8.
29. UKCCCR. UKCCCR guidelines for the use of cell lines in cancer research. Br J Cancer. 2000;82(9):1495–1509. doi:10.1054/bjoc.1999.1169.
30. Geraghty RJ, Capes-Davis A, Davis JM, Downward J, Freshney RI, Knezevic I, Lovell-Badge R, Masters JRW, Meredith J, Stacey GN, et al. Guidelines for the use of cell lines in biomedical research. Br J Cancer. 2014;111(6):1021–1046. doi:10.1038/bjc.2014.166.
31. Reid YA, As D, Steebbergen K, Hay RJ. Cell line methods. Encyclopedia of Life Sciences. 2001. doi:10.1038/npg.els.0002559.
32. Drexler HG, Dirks WG, Matsuo Y, MacLeod RAF. False leukemia-lymphoma cell lines: an update on over 500 cell lines. Leukemia. 2003;17(2):416–426. doi:10.1038/sj.leu.2402799.
33. Schweppre R, Kloppe J, Korch C, Pugazhenthhi U, Benezza M, Knauf JA, Fagin JA, Marlow LA, Copland JA, Smalldridge RC, et al. Deoxyribonucleic acid profiling analysis of 40 human thyroid cancer cell lines reveals
cross-contamination resulting in cell line redundancy and misidentification. J Clin Endocrinol Metab. 2008;93 (11):4331–4341. doi:10.1210/jc.2008-1102.

34. Phuchareon J, Ohta Y, Woo JM, Eisiele DW, Tetsu O. Genetic profiling reveals cross-contamination and misidentification of 6 adenoid cystic carcinoma cell lines: ACC2, ACC3, ACCM, ACCNS, ACCS and CACC. J Natl Cancer Inst. 2010;102(4):271–274. doi:10.1093/jnci/djp099.

35. Zhao M, Sano D, Pickering CR, Jasser SA, Henderson YC, Clayman GL, Sturgis EM, Ow TJ, Lotan R, Carey TE, et al. Assembly and initial characterization of a panel of 85 genomically validated cell lines from diverse head and neck tumor sites. Clinical Cancer Research. 2011;17(13):7248–7264. doi:10.1158/1078-0432.CCR-11-0690.

36. Moher D, Liberati A, Tetzlaff J, Altman G. Preferred reporting for systematic review and meta-analyses: the PRISMA statement. BMJ. 2009;339:b2535. doi:10.1136/bmj.b2535.

37. Pozos AMG, Garrocho JAR, Moreno UF, y Pérez MP. Capítulo 2. Búsqueda apropiada de la información (pp. 11–26). In: Moscoso AA, editor. Manual de Investigación Clínica. México, DF: Manual Moderno. 2012.

38. Auersperg N, Hawryluk AP. Chromosome observations on three epithelial-cell cultures derived from carcinomas of the human cervix. J Nat Cancer Inst. 1962;28:605–627.

39. Auersperg N. Long-Term cultivation of hypodiploid human tumor cells. J Nat Cancer Inst. 1964;32:135–163.

40. Auersperg N. Histogenetic behavior of tumors. III. Possible relationships to patterns of glycolysis. J Nat Cancer Inst. 1972;48:1589–1596.

41. Friedl F, Kimura I, Osato T, Ito Y. Studies on a new human cell line (sila) derived from carcinoma of uterus. I. Its establishment and morphology. Exp Biol Med. 1970;135(2):543–545. doi:10.3181/00379727-135-35091a.

42. Freedman R, Bowen J, Leibovitz A, Pathak S, Siciliano M, Gallager H, Giovannelli B. Characterization of a cell line (SW756) derived from a human squamous carcinoma of the uterine cervix. Vitro. 1982;18(8):719–726. doi:10.1016/BF00796428.

43. Popescu NC, Amsbaugh SC, DiPaolo JA. Human papillomavirus type 18 DNA is integrated at a single chromosome site in cervical carcinoma cell line SW756. J Virology. 1987;51(5):1682–1685. doi:10.1128/JVI.51.5.1682-1685.1987.

44. Nozawa S, Udagawa Y, Ontha H, Kurihara S, Fishman W. Newly established human uterine cervical cancer cell line (SKG-III) with Ragan isoezmonic, human chorionic gonadotropin beta-subunit, and pregnancy-specific beta 1-glycoprotein phenotypes. Cancer Res. 1983;43:1748–1760.

45. Shirasawa H, Tomita Y, Sekiya S, Takamizawa H, Simizu B. Integration and transcription of human papillomavirus type 16 and 18 sequences in cells line derived from cervical carcinomas. J Gen Virol. 1987;68(2):583–591. doi:10.1099/0022-1317-68-2-583.

46. Kelland LR, Burgess L, Gordon S. Characterization of four new cell lines derived from human squamous carcinomas of the uterine cervix. Cancer Res. 1987;47:4947–4952.

47. Spence RP, Murray A, Banks L, Kelland LR, Crawford L. Analysis of human papillomavirus sequences in cell lines recently derived from cervical cancers. Cancer Res. 1988;48:324–328.

48. Han X, Lyle R, Eustace DLS, Jewers RJ, Parrington JM, Das A, Chana T, Dagg B, Money S, Bates TD, et al. XH1 – a new cervical carcinoma cell line and xenograft model of tumour invasion, metastasis and regression. Br J Cancer. 1991;64(4):645–654. doi:10.1038/bjc.1991.376.

49. Monroy AG, Rangel RC, Trejo CB, Ramirez JG, Martinez R, Weiss BS. Establecimiento de siete estipres celulares provenientes de biopsias de cérvix normal y con cáncer cervical-uterino y sus diferentes contenidos y localizaciones de desmogéina-1. Rev Mex Oncol. 1992;7:69–76.

50. Kim JW, Cho YH, Lyu MS, Lee CG, Jung JK, Kim SJ, Namkoong SE. Establishment and characterization of a cell line (CUMC-3) derived from a human squamous carcinoma of the uterine cervix. Gynecol Oncol. 1995;57(1):47–60. doi:10.1006/gyno.1995.1098.

51. Kim JW, Lee CG, Cho YH, Kim JH, Kim SJ, Kim HK, Park TC, Song SK, Namkoong SE. CUMC-6, a new diploid human cell line derived from a squamous carcinoma of the uterine cervix. Gynecol Oncol. 1996;62(2):230–240. doi:10.1006/gyno.1996.0221.

52. Braun L, Mikumo R, Mark HF, Lauchlan S. Analysis of the growth properties and physical state of the human papillomavirus type 16. Am J Pathol. 1993;143:833–844.

53. Mark HF, Hane R, Mikumo R, Lauchlan S, Beauregard L, Braun L. Cytogenetic characterization of three cell lines derived from primary cervical tumors. Ann Clin Lab Sci. 1995;25:185–199.

54. Sonoda K, Nakashima M, Saito T, Amada S, Kamura T, Nakano H, Watanabe T. Establishment of a new human cervical adenocarcinoma cell-line, siso, and its reactivity to anticancer reagents. Int J Oncol. 1995;6(3):1099–1104. doi:10.3892/ijo.6.5.1099.

55. Chou CY, Chen YH, Tzeng CC, Cheng YC, Chang CF, Chen TM. Establishment and characterization of a human-papillomavirus negative, p53-mutation negative human cervical cancer cell line. Cancer Lett. 1996;102(1–2):173–181. doi:10.1016/0304-8853(96)04157-2.

56. Isaka K, Nishi H, Osaka Y, Miyata M, Hokamura M, Nakada T, et al. Establishment of a HPV and p53-mutation-negative human cell line (CA) derived from a squamous carcinoma of the uterine cervix. Gynecol Oncol. 2004;92(1):15–21. doi:10.1016/j.ygyno.2003.09.014.

57. Ishiwata I, Tokieda Y, Iguchi M, Tamagawa T, Ishiwata I, Kiguchi K, Ishikawa H. Establishment and characterization of human uterine cervical epithoderm carcinoma cell line HHUS containing HPV 59 DNA. Hum Cell. 2004;17(3):151–156. doi:10.1111/j.1749-0774.2004.tb00332.x.

58. Javed S, Sharma BK, Sood S, Sharma S, Bagga R, Bhattacharyya S, Rayat CS, Dhivali L, Srinivasan R. Significance of CD133 positive cells in four novel HPV-16 positive cervical cancer derived cell lines and biopies of invasive cervical cancer. BMC Cancer. 2018;18(1):357. doi:10.1186/s12888-018-4237-5.

59. Kiguchi K, Ishiwata I, Tokieda Y, Iguchi M, Ishiwata I, Iwata M, Ishizuka B, Yoshikawa H, Tachibana T, Hashimoto H, et al. Establishment and Characterization of a Lymphoepithelial-like Carcinoma Cell Line (HUVEC) Derived from the Human Uterine Cervix. Hum Cell. 2002;15(2):97–102. doi:10.1111/j.1749-0774.2002.tb00104.x.

60. United Nations Educational, Scientific and Cultural Organization. UNESCO science report, towards 2030: Executive summary. UNESCO publishing: France. 2015.

61. Wilbanks GD, Leipus E, Tsurumoto D. Chapter 2 tissue culture of the human uterine cervix. Normal Human Tissue and Cell Culture B. Endocrine, Urogenital, and Gastrointestinal Systems. 1980:29–50. doi:10.1001/0001-0769(0806)0677-9.

62. Masters JR, Stacey GN. Changing medium and passaging cell lines. Nat Protoc. 2007;2(9):2276–2284. doi:10.1038/nprot.2007.319.

63. Organización Mundial de la Salud. Principios rectores de la OMS sobre trasplante de células, tejidos y órganos humanos. 2010. Encontrado el 27/07/2020 en: https://www.who.int/transplantation/Guiding_PrinciplesTransplantation_WHA63.2232.pdf

64. International Cell Line Authentication Committee (ICLAC). Database of cross-contamination or misidentified cell lines. Versión. 2014;7(2).
66. Reid Y, Storts D, Riss T, et al. Authentication of human cell lines by STR DNA profiling analysis. In: Markosssian S, Sittamplam G and Grossman A, editors. Assay guidance manual. Bathesda (MD): Eli Lilly & Company and the National Center for Advancing Translational Sciences; 2013 (Internet).

67. Marx V. Cell-Line authentication demystified. Nat Methods. 2014;11(5):483–488. doi:10.1038/nmeth.2932.

68. Liaw K, Hildesheim A, Burk RD, Gravitt P, Wacholder S, Manos MM, Scott D, Sherman M, Kurman R, Glass A, et al. A prospective study of human papillomavirus (HPV) type 16 DNA detection by polymerase chain reaction and its association with acquisition and persistence of other HPV types. J Infect Dis. 2001;183(1):8–15. doi:10.1086/317638.

69. Cuschieri KS, Cubie HA, Whitley MW, Seagar AL, Arends MJ, Moore C, et al. Multiple high risk HPV infections are common in cervical neoplasia and young women in a cervical screening. J Clin Pathol. 2004;57(1):68–72. doi:10.1136/jcp.57.1.68.

70. Trottier H, Mahmud S, Costa MC, Sobrinho JP, Duarte-Franco E, Rohan TE, et al. Human papillomavirus infections with multiple types and risk of cervical neoplasia. Cancer Epidemiol Biomarkers Prev. 2006;15(7):1274–1280. doi:10.1158/1055-9965.EPI-06-0129.

71. Gargiulo F, De Francesco MA, Schreiber C, Caravolo G, Salinaro F, Valloncini B, Manca N. Prevalence and distribution of single and multiple HPV infections in cytologically abnormal cervical samples from Italian women. Virus Res. 2007;125(2):176–182. doi:10.1016/j.virusres.2006.12.017.

72. Spinillo A, Dal Bello B, Gardella B, Roccio M, Dacco MD, Silini EM. Multiple human papillomavirus infection and high grade cervical intraepithelial neoplasia among women with cytological diagnosis of atypical squamous cells of undetermined significance or low grade squamous intraepithelial lesions. Gynecol Oncol. 2009;113(1):115–119. doi:10.1016/j.ygyno.2008.12.037.

73. Pista A, Oliveira A, Verdasca N, Ribeiro F. Single and multiple human papillomavirus infections in cervical abnormalities in Portuguese woman. Clin Microbiol Infect. 2011;17(6):941–946. doi:10.1111/j.1469-0691.2010.03387.x.

74. Fogh J, Wright WC, Loveless JD. Absence of HeLa cell contamination in 169 cell lines derived from human tumors. J Natl Cancer Inst. 1977;58(2):209–305. doi:10.1093/jnci/58.2.209.

75. Yoshino K, Limura E, Saijo K, Iwase S, Fukami K, Ohno T, Obata Y, Nakamura Y. Essential role for gene profiling analysis in the authentication of human cell lines. Hum Cell. 2006;19(1):43–48. doi:10.1111/j.1749-0774.2005.00007.x.

76. Wenger SL, Senft JR, Sargent LM, Bamezai R, Bairwa N, Grant SG. Comparison of established cell lines at different passages by karyotype and comparative genomic hybridization. Biosci Rep. 2004;24(6):631–639. doi:10.1007/s10540-005-2797-5.

77. Rutzky CP, Kaye CJ, Siciliano CM, Kahan BD. Longitudinal karyotype and genetic signature analysis of cultures human colon adenocarcinoma cell lines LS180 and LS174T. Can Res. 1980;40:1443–1448.