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ORIGINAL RESEARCH

Genetic profiles of cervical tumors by high-throughput sequencing for personalized medical care

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Abstract

Cancer treatment is facing major evolution since the advent of targeted therapies. Building genetic profiles could predict sensitivity or resistance to these therapies and highlight disease-specific abnormalities, supporting personalized patient care. In the context of biomedical research and clinical diagnosis, our laboratory has developed an oncogenic panel comprised of 226 genes and a dedicated bioinformatic pipeline to explore somatic mutations in cervical carcinomas, using high-throughput sequencing. Twenty-nine tumors were sequenced for exons within 226 genes. The automated pipeline used includes a database and a filtration system dedicated to identifying mutations of interest and excluding false positive and germline mutations. One-hundred and seventy-six total mutational events were found among the 29 tumors. Our cervical tumor mutational landscape shows that most mutations are found in PIK3CA (E545K, E542K) and KRAS (G12D, G13D) and others in FBXW7 (R465C, R505G, R479Q). Mutations have also been found in ALK (V1149L, A1266T) and EGFR (T259M). These results showed that 48% of patients display at least one deleterious mutation in genes that have been already targeted by the Food and Drug Administration approved therapies. Considering deleterious mutations, 59% of patients could be eligible for clinical trials. Sequencing hundreds of genes in a clinical context has become feasible, in terms of time and cost. In the near future, such an analysis could be a part of a battery of examinations along the diagnosis and treatment of cancer, helping to detect sensitivity or resistance to targeted therapies and allow advancements towards personalized oncology.

Introduction

Cancer treatment faces a major evolution since the advent of targeted therapies. It is now possible to specifically treat tumors presenting with a clearly identified genetic alteration(s). It is the case of nonsmall-cell lung cancers associated with *EGFR* mutations that responds to specific tyrosine kinase inhibitors [1]. This type of therapy can

only be effective in the case of a known molecular target. It requires the characterization of various alterations (commonly called actionable mutations) that a tumor may accumulate. Establishment of such genetic profiles would allow sensitivity, resistance, and toxicity predictions for such therapies [2]. Sequences from many human genomes have shown a wide interindividual genetic heterogeneity [3]. Variations among individuals are attributed to

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"germline" mutations and constitute the individual's inherited genetic characteristics. These mutations, however, should be distinguished from other mutations that may occur during the individual's life, and also, might only be observed in one organ or some tissue. These events are called somatic mutations and have been shown to provoke various oncogenic processes [2] (they are called "driver" mutations).

Many research teams, brought together in international consortiums, are engaged in the characterization of mutations causing tumorigenesis. The Catalog Of Somatic Mutations In Cancer (COSMIC) Project [4], for example, aims at creating a database gathering all somatic mutations already described. The International Cancer Genome Consortium plans to build a comprehensive catalog of somatic mutations in tumors of 50 different types/subtypes of cancer, with additional epigenomic and transcriptomic information, with the aim of highlighting differences and common abnormalities across tumor types [5]. These works on tumor genetic profiling have demonstrated that tumors of different types can share protumorigenic signaling pathways, defined by common driver mutations which could also be actionable [6]. Consequently, it makes sense to explore the presence or absence of mutations within these driver genes, to identify biomarkers for sensibility or resistance to treatments, and to indicate those patients most likely to benefit from targeted therapies.

The number of genes to test, in order to establish a cancer gene map, is huge [7]. This approach is made possible by a new generation of sequencing devices (Next Generation Sequencing, NGS), able to analyze the equivalent of several entire genomes [8]. However, sequencing a whole genome can be quite long, regardless of analysis time for such an extensive quantity of data. Another approach consists of limiting the analysis to exomes or several hundreds of genes, using targeted methods. Using a rational approach, which targets a panel consisting of a few hundred genes, considered to be "actionable" and/ or "driver", it is possible to sequence and characterize a tumor, depending on its major molecular characteristics. The main advantages of this approach lie in the reduced delay for reporting the results and limited costs, compatible with a diagnostic use, while maintaining sufficient sequencing quality to detect somatic variations [9].

The sensitivity of these methods depends on bioinformatic tools specifically developed to highlight somatic mutations in cancer, by comparing healthy tissue and tumor DNA from the same individual (paired-mode). These tools verify that a somatic mutation found in tumor does not actually correspond to germline mutation or systematic false-positive mutation, found in healthy tissues [10]. In some cases, it is difficult to obtain the matched

healthy tissue in a clinical context, depending on legislative (local or national), ethical or logistical considerations; a constraint that must be overcome. On the other hand, the detection of mutations strongly depends on algorithms and their adjusted parameters used. In addition, their sensitivity depends on the type and rate of mutation present in the tumor. To date, there is no global consensus for the use of a particular algorithm or mode of analysis [10].

We report on the genetic profiles of cervix uteri tumors which represents the fourth leading cause of death by cancer in females worldwide [11]. It is well-known that some human papillomavirus (HPV) types are the cause of this cancer and its development is linked to the stable insertion of the HPV genome into the tumor's DNA [12]. Cancer of cervix uteri can go undetectable for years and years, and is associated with a 5-year survival rate of 68% [13], depending on clinical stage at presentation: the 5-year relative survival rate is 91%, 58%, and 17% for patients with localized, regional, and metastatic disease, respectively (http://seer.cancer.gov/statfacts/html/cervix. html). Moreover, up to one-third of patients will develop recurrent tumors mostly within 2 years after initial treatments [14]. Most recurrences being detected by imaging or medical examination, it is essential that highly specific, sensitive and less invasive markers are identified to predict response to treatment, disease progression, and to develop personalized therapies.

In our laboratory, we have developed the sequencing of a panel of genes associated with tumors of cervix uteri, including exons within 226 genes chosen for their actionable nature and their potential implication in cervical tumorigenesis (Table 1). To overcome the lack of matched healthy tissue, an average healthy tissue has been generated in silico from nonmatched healthy tissues at our disposal. The aim of the study was to prove that a tumor can be managed in a clinical context using NGS technology and automated bioinformatic pipeline and to identify potential actionable mutations that could have a direct link with treatments.

Material and Methods

Tumor samples

Twenty-five squamous cell carcinoma and four adenocarcinoma tumor samples were obtained under approved protocols from the Curie Institute in Paris, France (Table 2). Seven healthy tissue samples were obtained from cervical surgical specimen provided by the Anatomic Pathology laboratory of cancer center François Baclesse in Caen, France. All patients gave written informed consent before entering the study.

Table 1. Genes selected for cervix uteri cancer panel.

ABL1	CDK8	FGFR3	LTK	NSD1	SMAD2
ABL2	CDKN1A	FGFR4	MAP2K1	NTRK1	SMAD3
AFF3	CDKN2A	FHIT	MAP2K2	NTRK2	SMAD4
AKT1	CEACAM5	FKBP9	MAP2K4	NTRK3	SMARCA4
AKT2	CEBPA	FLT1	MCL1	PAK3	SMARCB1
AKT3	CHEK1	FLT3	MDM2	PARP1	SMO
ALK	CHEK2	FLT4	MDM4	PAX5	SOCS1
APC	CREBBP	FOXP4	MED1	PDGFRA	SOX10
AR	CRKL	GATA1	MEN1	PDGFRB	SOX2
ARFRP1	CROCC	GLIS2	MET	PDPK1	SRC
ARID4A	CSF1R	GNA11	MGMT	PIK3CA	STK11
ATM	CSMD1	GNAQ	MITF	PIK3R1	SUFU
AURKA	CTLA4	GNAS	MLH1	PKHD1	TBX22
AURKB	CTNNB1	GPR124	MLL	PLCG1	TCF4
BCL2	CTTN	GUCY1A2	MLL2	PLEKHO2	TENM1
BCL2A1	DAPK1	HIC1	MLL3	PRDM9	TERT
BCL2L1	DDX3X	HNF1A	MPL	PRKDC	TET2
BCL2L2	E2F3	HOXA3	MRE11A	PTCH1	TGFBR2
BCL6	EGFR	HRAS	MSH2	PTEN	TIMP2
BRAF	EMSY	HSP90AA1	MSH6	PTK2	TIMP3
BRCA1	EPHA3	IDH1	MTOR	PTK2B	TNFAIP3
BRCA2	EPHA5	IDH2	MUC1	PTPN11	TNFRSF10C
CADM1	EPHA6	IGF1R	MYB	PTPRD	TNFRSF10D
CASP8	EPHA7	IGF2R	MYC	RAD51	TOP1
CASZ1	EPHB1	IKBKE	MYCL1	RAD51B	TP53
CAV1	EPHB4	IKZF1	MYCN	RAF1	TP63
CBL	EPHB6	INHBA	MYH15	RARA	TP73
CCND1	ERBB2	IRF6	NF1	RARB	TSC1
CCND2	ERBB3	IRS2	NF2	RASSF1	TSC2
CCNE1	ERBB4	JAK2	NFE2L2	RB1	TSHR
CCNL1	ESR1	JAK3	NFIB	REL	USP9X
CDC73	EZH2	KDR	NKX2-1	RET	VHL
CDH1	FAM123B	KEAP1	NOTCH1	RICTOR	WT1
CDH2	FANCC	KIAA0774	NOTCH2	ROS1	ZBTB7C
CDH20	FANCF	KIT	NOTCH3	RPS6KB1	ZNF668
CDH5	FBXW7	KLF5	NOTCH4	RPTOR	ZNF91
CDK4	FGFR1	KRAS	NPM1	RUNX1	
CDK6	FGFR2	LRP1B	NRAS	RUNX1T1	

Sample preparation and next-generation sequencing

Sequencing methods were described previously [15], enabling analysis of two tumors and one healthy tissue sample in the same sequencing run. The SureSelect-targeted enrichment process (Agilent, Santa Clara, CA) was performed after combining the indexed samples, equimolarly. Libraries were then sequenced on a MiSeq (Illumina, San Diego, CA), using the paired-end 2×150 bp method (full protocol available on request).

Bioinformatic analysis

The CASAVA suite v1.8 (Illumina) was used for demutiplexing and generating fastq files. These raw data were then analyzed by a home-made pipeline (BAPT, Bioinformatic

Analysis Pipeline Toolkit), to automate alignment, variantcalling, and annotation steps. Reads were first mapped to a reference genome (hg19) using BWA algorithm [16] and output files were reworked following GATK [17] (Genome Analysis ToolKit, Broad Institute) best practices, especially to calculate quality scores and undergo local realignments. Five variant-callers were used to call single-nucleotide substitutions and insertion-deletion (indels) events: HaplotypeCaller [18], UnifiedGenotyper [18], LofreqStar [19], Varscan2 [20], and MuTect [21]. The BAM files from seven healthy samples were used by randomly selecting reads from each file, to create an average healthy synthetic file (using SAMtools [22]). This "synthetic" BAM file was used in paired-mode during the variant-calling step with the programs LofregStar, MuTect, and VarScan2. All variants found by the variant callers were annotated with Alamut-Batch (Interactive Biosoftware, Rouen, France) and integrated into a database called CanDiD

Table 2. Demographic, histological, and biological characteristics.

Patient no.	Histological type	Tumor stage (FIGO)	Age at diagnosis	HPV type		
1	SCC	IIB	40	HPV 18		
2	SCC	IB	33	HPV 18		
3	Adenocarcinoma	II	45	HPV 18		
4	SCC	IB	37	HPV 18		
5	SCC	IB	34	HPV 16		
6	SCC	IB2	47	HPV 16		
7	Adenocarcinoma	IIB	49	HPV 16		
8	SCC	IVB	57	HPV 16		
9	SCC	IB1	33	HPV 16		
10	SCC	NA	NA	HPV45		
11	SCC	IIB	60	HPV 16		
12	SCC	IIB	42	HPV 18		
13	SCC	IIB	47	HPV 16, 18		
14	SCC	IIB	34	HPV 16		
15	SCC	IIB	68	HPV 16		
16	SCC	IIB	43	HPV 18		
17	SCC	IIB	44	HPV 51		
18	SCC	IIB	65	HPV 33		
19	SCC	IIB	45	HPV 73		
20	SCC	IIIB	43	HPV 16		
21	SCC	IIB	53	HPV 16		
22	Adenocarcinoma	IB2	42	HPV 16		
23	Adenocarcinoma	IIB	54	HPV-		
24	SCC	IIB	54	HPV 18		
25	SCC	III	33	HPV 16		
26	SCC	IIB	55	HPV 16		
27	SCC	IV +	44	HPV 73		
	metastasis					
28	SCC	IB1	25	HPV 18		
29	SCC	IB2	31	HPV-		

FIGO, International Federation of Gynecology; HPV, human papillomavirus; SCC, squamous cell carcinoma; NA, not available.

(Cancer Diagnostic Database, PostgreSQL) (Fig. S1). Variants were retrieved from CanDiD database according to five criteria: (1) the variant was found on a list of preferred transcripts; (2) the variant was in targeted zones; (3) the variant was within the coding sequence or ±10 bp within the intronic sequences; (4) the variant was in a canonical splicing mutation sites or when outside they induced a 15% decrease of MaxEntScan score and a 5% decrease of the SpliceSiteFinder score [23] (hereafter called splicing mutation); and (5) the variant was observed 10 times at most in the Exome Sequencing Project Database (ESP; http://evs. gs.washington.edu.EVS/). Identified variants passed through a home-made filtration system (scripted on Python programming language, available on request) which compares sequencing data from the five variant-callers and tumor samples versus healthy samples, in order to rule out false-positive and germline variants. Data from each variant were first compared to both tumor and healthy tissue samples, to evaluate whether the variant have a greater presence in the tumor than in the healthy samples. Variants were then filtered based on their quality score (the main criteria being the PHRED score and depth of coverage). The main objective was to obtain, for each tumor, a list of variants with a high probability of being somatic. Only variants seen by at least 2 variant callers were further considered.

Results were annotated and interpreted in sequence with: (1) two mutation impact prediction systems SIFT [24], Polyphen [25], (2) InterPro Database [26], (3) Clinical Trials.gov Database [27], (4) dbSNP [28], (5) COSMIC [4], (6) TARGET [29].

In the present study, all genes that constitute a target for the US Food and Drug Administration-approved targeted therapies were called actionable genes, regardless of the type of cancer. Similarly, a missense mutation was classified as deleterious if scored as "deleterious" by SIFT algorithm or "Probably Damaging" by Polyphen algorithm. Inactivating mutations (PTC, premature codon termination and splicing mutations) were directly considered as deleterious.

Results

Tumor samples from 29 patients with cervical cancer (stade I B to IV B), including 25 squamous cell carcinomas and 4 adenocarcinomas, have been sequenced (Table 2), as well as seven samples from healthy cervical biopsy ("control" samples). The time from DNA extraction to acquisition of most likely somatic mutations was about height working days (Fig. 1). The sequencing process produced an average of 8,078,023 reads per sample, with an average sequencing depth of 268× and >92% of nucleotides covered ≥50-fold. Consequently, only variants with an allele ratio of more than 5% were called. After data processing in the BAPT pipeline, 11,267 variants were included in the CanDiD database. After extraction from the database, 2746 missense mutations were collected from the various variant callers (Fig. 2A). After passing through the filtration system, only 220 mutations were retained (Fig. 2B; Table S2). Most of the mutations deleted by the filtration system were germline mutations detected by HaplotypeCaller or UnifiedGenotyper (from the control samples) or false-positive selected by Varscan due to its lack of specificity. Among all mutations found in the 29 tumor samples, 41% (91/220) were identified by all variant callers, assuring accuracy of the mutations detected. Only variants detected by two or more variantcallers were selected leading to 156 missense mutations selected, among which 29 are already referenced in the COSMIC database. Seven nonsense mutations, nine small insertion/deletions (indels) inducing PTC and four splicing mutations were also identified. Altogether, they represent an average of 6.1 mutations per tumor sample. For each patient, 75 percent of mutations were considered

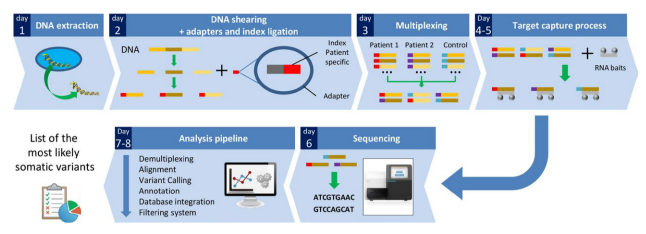


Figure 1. Sequencing workflow's major steps. After DNA extraction (Day 1), DNA molecules are sheared by sonification, and DNA fragments are ligated to adapters containing a patient-specific index (Day 2). DNA from 2 patients and 1 control are pooled equimolarly together during the multiplexing step (Day 3). Regions of interest (exons from 226 genes) are retrieved by a targeted enrichment system with biotinylated baits (Day 4–5). Then DNA is sequenced in an Illumina MiSeq (Day 6) and sequencing raw data are processed by the bioinformatic pipeline (Day 7–8), to extract the most likely somatic variations.

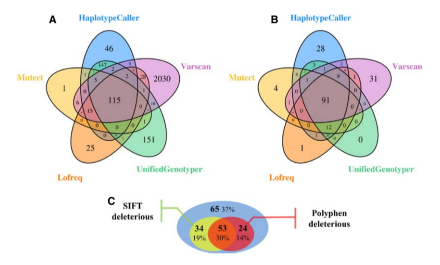


Figure 2. Representation of missense mutations found by each variant caller. (A) Mutations extracted directly from CANDID database (total: 2746). (B) mutations remaining after passing through filtration system (total: 221). (C) Proportion of mutations detected by at least 2 variant-callers classified as deleterious by SIFT or POLYPHEN.

deleterious (by SIFT or POLYPHEN algorithm) (Fig. 2C, Table S2).

No somatic missense mutation was found in the tumor suppressor gene TP53, which is consistent with molecular characteristics of cervix uteri tumors [30]. Similarly, missense mutations were found in the oncogene PIK3CA (n=8), the KRAS gene implicated into anti-EGFR therapies resistance (n=4) and in the FBXW7 gene, recently described in the cervix uteri tumors mutational landscape (n=3) (Table S2). Other missense mutations were found in tumor suppressor genes, including the susceptibility gene for breast cancer BRCA2 (n=6) or the RAS family negative regulator NF1 (n=1). Twenty inactivating

mutations (PTC and splicing mutations) were also identified in tumor suppressor genes including *RB1* (n = 1), *NF1* (n = 3) or *MLH1* (n = 2) (Table 3).

Among other actionable genes, deleterious missense mutations were observed in *ALK* (n=4), *AR* (n=1), *EGFR* (n=3), *ERBB2* (n=1), *MET* (n=1), *mTOR* (n=2), *PDGFRA* (n=1) and *RAF1* (n=2) (Table 4). Considering only deleterious mutations, 48% (n=14/29) had at least one mutation in an actionable gene (Fig. 3A and B).

Several clinical trials (www.clinicaltrials.gov database) are looking for mutations in specific genes to evaluate sensitivity or resistance to approved, or in clinical trials, targeted therapies, among which PIK3CA, ALK, EGFR,

Table 3. Inactivating mutations.

Coding effect	Gene	Coding DNA sequence ¹	Protein sequence ¹	Transcript	Patients tumor sample no.
Splicing mutation	CBL	c.1096–1G>A	p. ?	NM_005188	5
Potential splicing	mTOR	c.2514+3G>C	p. ?	NM_004958	15
mutations	MYH15	c.1993-14G>A	p. ?	NM_014981	14
	NF1	c.1392+5G>T	p. ?	NM_001042492	21
Nonsense mutations	C11orf30	c.3004C>T	p.Gln1002*	NM_020193	26
	CEACAM5	c.646G>T	p.Glu216*	NM_004363	26
	EPHA5	c.2458G>T	p.Gly820*	NM_004439	6
	FBXW7	c.1053G>A	p.Trp351*	NM_033632	26
	MLH1	c.1630C>T	p.Gln544*	NM_000249	20
		c.755C>T	p.Ser252*		5
	RB1	c.1399C>T	p.Arg467*	NM_000321	5
Frameshift mutations	CASP8	c.790dup	p.Val264Glyfs*13	NM_001080125	5
	CHEK2	c.1229del	p.Thr410Metfs*15	NM_001005735	11
	CREBBP	c.4477dup	p.Ile1493Asnfs*26	NM_004380	5
	FGFR2	c.962dup	p.Asn321Lysfs*21	NM_022970	5
	FGFR4	c.2396_2403del	p.Gly799Aspfs*12	NM_213647	7
	NF1	c.5907_5908del	p.Arg1970Serfs*6	NM_001042492	5
		c.2033del	p.Pro678Argfs*10		
	NF2	c.301del	p.Tyr101llefs*22	NM_181832	16
	NOTCH2	c.6909del	p.lle2304Leufs*2	NM_024408	5

Mutations are classified by mutation type.

Table 4. Deleterious mutations found in actionable genes.

Gene	Drugs in relation with gene of interest	Mutation	Number of tumors	Transcript	Associated clinical trial
ALK	Crizotinib, Ceritinib	A1266T	4	NM_004304	NCT01548144,
		A1234V			NCT01744652
		V1149L			
		R1120Q			
4R	Entuzalamide, Abiraterone	K809N	1	NM_000044	_
GFR	Cetuximab, Panitumumab, Erlotinib,	S511Y	3	NM_005228	NCT00770263
	Gefetinib, Afatinib, Vandetanib	T259M			
		A611T			
RBB2	Trastuzumab, Pertuzumab, Lapatinib	L696F	1	NM_001005862	NCT01953926
KRAS	Resistance to Cetuximab and others	G12D	4	NM_033360	_
		G13D			
ЛЕТ	Cabozantinib	L342F	1	NM_001127500	_
nTOR	Temsirolimus, Everolimus	M813I	1	NM_004958	_
DGFRA	Regorafenib	P441L	1	NM_006206	NCT02029001
RAF1	Regorafenib	C96F	2	NM_002880	_
		Q255H			

Genes are linked to targeted therapies already approved by Food and Drug Administration in at least one indication. All potential drug targets of each therapy are considered.

mTOR, PTCH1, SMO, and PIK3R1 are included (Table S2). Overall, 59% of patients (17/29) would be eligible for potential inclusion in a clinical trial (Fig. 3A and B), based on carrier gene mutations and tumor origin.

Discussion

The aim of the study was to demonstrate in a clinical context the feasibility of detecting somatic variations, in

cervical cancers, using a targeted sequencing approach. The use of a "synthetic healthy tissue sample" (see above Material and Methods section) from the sequencing data of several unpaired healthy tissue samples allows the use of somatic variant-callers (which compares healthy tissues from tumors to extract somatic mutations) to detect poorly represented mutations. This approach, however, probably tolerates a few germline mutations which cannot be differentiated from somatic mutations. In our study, some

¹Nomenclature according HGVS guidelines (Human Genome Variation Society).

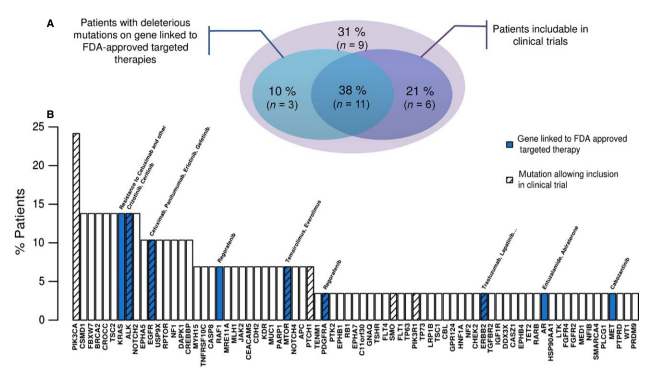


Figure 3. Distribution of deleterious mutations among the 29 tumors. A gene is considered actionable if linked to targeted therapies approved by Food and Drug Administration. A mutation is classified as deleterious if considered as such by SIFT algorithm or POLYPHEN algorithm. (A) Proportion of tumors with a deleterious mutation on gene considered actionable (blue) and proportion of patients for whom their deleterious mutations could allow inclusion in a clinical trial (purple). (B) Representation of most mutated genes across tumors.

mutations found in BRCA2 (hereditary predisposition for breast and ovarian cancer) or in MLH1 (hereditary predisposition to Lynch syndrome) genes through the genetic characterization of tumors could be germline (Table S2). The identification of germline mutations in genes in relation with mendelian diseases must be taken into account and the appropriate ethical issues should be considered, a subject still of debate. Tools and methods developed in our laboratory should not be considered as an automatic interpretation system; they should be considered as help to ease work discerning the most relevant mutations. For instance, with our data, the number of mutations to analyze drops from about 388 to 6 for one tumor, which represents a huge time-saving for biologists who interpret data in a clinical context. Moreover, analysis time (±8 working days) is consistent with clinical practice, and should improve patient healthcare in a comprehensive way.

No somatic alterations were detected in the tumor suppressor gene *TP53* which seems consistent throughout our model. In cervical carcinomas *TP53* inactivation is linked to virus integration, which results in the expression of the E6 oncoprotein, able to induce the *TP53* degradation via the ubiquitin pathway [30]. Previous studies have already identified genes frequently mutated

in cervical carcinomas, such as PIK3CA [31], KRAS [32] (the most mutated genes in cervix uteri cancer according to COSMIC data) or FBXW7 [33]. PIK3CA E545K (n = 4)and E542K (n = 2) have strong oncogenic properties due to increased kinase activity [34], consequences in growth factor-independent cell proliferation, resistance to apoptosis, and increased invasion and cell migration. This gene represents a prime target for drug development, and even if there is no available approved treatment, several clinical trials are currently testing PIK3CA selective inhibitors (ClinicalTrials ID: NCT01708161; NCT01928459). Two major mutations were also found in KRAS, G12D (n = 1) and G13D (n = 3), well known in colorectal cancer to predict a resistance to EGFR antibodies [35]. FBXW7 is a gene recently described as mutated in cervical carcinoma [33] with two missense mutations already described, R505G and R456C. There was also a novel mutation at a position already found mutated, R479Q. All of these point mutations are located in the WD repeats domain. This domain is used to form a complex in charge of degrading several products of proto-oncogenes (including MYC, CCNE1, NOTCH1, and JUN [36]. We found a novel truncating mutation W351*; it occurs just upstream of the WD domain (uni-Prot ID: Q969H0). Therefore, pipeline implementation

seems consistent, since key elements form cervical tumorigenesis were found.

Considering all of the actionable genes as defined before (Table 4), deleterious mutations could influence sensitivity or resistance to treatments. ALK mutations V1149L and A1266T are located next to previously described mutations L1152R [37] and G1269A [38], which have been identified as resistance mutations to Crizotinib and other tyrosine kinase inhibitors, and could then have the same impact on treatment response. Another mutation (T259M) has been observed in a specific subpart of extracellular domain of EGFR, called the dimerization loop [39]. This loop is essential for receptor dimerization and hence activation. A previous study showed that mutations in this subpart were able to increase receptor oncogenic activity [40]. MET had also a new mutation at residue L342, which correspond to a specific region of SEMA domain, considered as MET ligand-binding domain [41] and useful to receptor dimerization and activation [42]. Other mutations classified as deleterious were detected in mTOR (M813I, splice mutation c.2514+3G>C) and PDGFRA (P441L) but their real impact on protein functionality or the effectiveness of related targeted therapies is still unknown.

Eighteen deleterious mutations identified from our cases reside in actionable genes (Table 3), although their impact is still unclear for many. Therefore, functional studies should be performed to identify potential drug sensitivities or resistances to these targets. Deleterious classification (SIFT and POLYPHEN algorithm) should be used with caution since, for instance, *PIK3CA* mutations E545K and E542K are classified as tolerated by SIFT despite the fact that they are known driver mutations.

Among driver genes investigated in our study, interesting mutations were found on *BRCA2*, with 2 variants of unknown signification, classified here as deleterious missense mutations (L2721R and A2770D) in the OB1 (oligonucleotide/oligosaccharide-binding 1) domain, which serves to bind single-strand DNA. Functional assays showed that mutations in this domain can be disease-causing [43]. Another mutation (V2066I) was observed in BRC repeat domains, essential for binding to RAD51 protein [44]. The new PARP inhibitor Olaparib, now available, could potentially be of interest in these BRCA mutated tumors. Unfortunately, for the moment this drug is mainly tested in breast and ovarian cancers. Altogether, the real new challenge for a tumor's genetic profiling is the correct interpretation of mutation.

Sixty-nine percent of patients could have had their treatment decision guided by the targeted sequencing of 226 genes, either by revealing sensitivity or resistance to available drugs or by directing patients toward treatment protocols or diagnostic procedures being tested (Fig. 3A).

Even though no therapeutic answer could be provided by our "oncopanel" for certain patients, they were all analyzed for HPV insertion status. The HPV insertion site constitutes another biomarker in cis, which can provide information for targeted therapies. HPV has therefore been found inserted into MYC, KLF5, KLF12, as well as RB1, and many other important and potentially actionable genes (A. Holmes, unpubl. data). Such tests might participate in the overall study of cancer biomarkers in both prognostic (assessing tumor outcomes and deciding whether or not to undergo treatment) and predictive (evaluating sensitivity and resistance to treatments to choose the most effective drug) fields [45]. The outcome of such studies strives toward personalized oncology, which is to provide the most appropriate treatment to the patient with the correct dosage. The overall benefits first and foremost concern patients, who receive the most effective and least toxic treatments. Such an approach in patient care has economic advantages too. The correct administration of the adequate drug instead of using other ineffective and expensive therapies appears to be cost-effective. It has been calculated that US health-care system could save up to \$740 million dollars by only treating patients with Cetuximab with a wild-type KRAS status [46]. Other studies suggest that pharmacogenomics can be a good way to improve cost-effectiveness [47]. However, more studies are needed in this field to assess the real economic impact of these therapies [48].

Interpretation of somatic mutations remains challenging because, except for a few mutations already described, the impact of each mutation requires functional studies. In some countries, the inability to obtain matched healthy tissue in a daily clinical practice makes detection of somatic mutations more difficult. On the other hand, sequencing a limited number of genes selected for their implication in tumor development is now feasible in a clinical setting (in term of costs and time), and the "oncopanel" approach allows a larger screening power for diagnosis or treatment orientation by identification of sensitivity or resistance to drugs.

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Conflict of Interest

None declared.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Figure S1. Somatic mutations analysis pipeline: BAPT.

Table S2. Somatic mutations found across the 29 patients.