A technical application of quantitative next generation sequencing for chimerism evaluation

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Abstract. At present, the most common genetic diagnostic method for chimerism evaluation following hematopoietic stem cell transplantation is microsatellite analysis by capillary electrophoresis. The main objective was to establish, through repeated analysis over time, if a complete chimerism was present, or if the mixed chimerism was stable, increasing or decreasing over time. Considering the recent introduction of next generation sequencing (NGS) in clinical diagnostics, a detailed study evaluating an NGS protocol was conducted, coupled with a custom bioinformatics pipeline, for chimerism quantification. Based on the technology of Ion AmpliSeq, a 44-amplicon custom chimerism panel was designed, and a custom bioinformatics pipeline dedicated to the genotyping and quantification of NGS data was coded. The custom chimerism panel allowed identification of an average of 16 informative recipient alleles. The limit of detection of the protocol was fixed at 1% due to the NGS background (<1%). The protocol followed the standard Ion AmpliSeq library preparation and Ion Torrent Personal Genome Machine guidelines. Overall, the present study added to the scientific literature, identifying novel technical details for a possible future application of NGS for chimerism quantification.

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Introduction

Allogenic hematopoietic stem cell transplantation (HSCT) is the predominant treatment used to cure malignant and non-malignant hematological disorders. The number of HSCTs conducted has increased due to an overall improvement in the safety of the procedure resulting from reduced-intensity conditioning regimens, and the availability of new donor sources, available in the national registries (1,2).

Reduced-intensity conditioning is widely used to avoid the complications of myeloablative conditioning to prepare for HSCT in the adult population. However, this procedure is associated with a high risk of complications that may result in graft loss. To prevent this, it is important to monitor chimerism for early intervention (3). Evaluation of chimerism status at regular intervals is useful to prevent risk of early graft rejection and relapse in patients suffering from malignant diseases. Quantification of the chimerism percentage is also a potential marker of minimal residual disease (MRD) for patients without suitable MRD markers and it provides useful information on graft vs. host disease and graft vs. tumor effects (4-6).

Chimerism analysis is a tool that allows to determination of the genotypic origin of post-transplantation hematopoiesis. Subsequent to HSCT, a patient presenting with 100.0% donor-origin cells during follow up is considered to have the status of complete chimerism (CC), patients in which the donor- and recipient-origin cells coexist have the status of mixed chimerism (MC) (7). Informative genetic markers are used to discriminate between recipient and donor genomes in order to detect the chimerism status (8).

At present, different approaches based on polymerase chain reaction (PCR) amplification of polymorphic DNA sequences (short tandem repeat, STR; single nucleotide polymorphism, SNP; and insertion/deletion, INDEL) are used for chimerism analysis. In the vast majority of laboratories, semi-quantitative fluorescent PCR of STRs is the procedure of choice for diagnostic purposes. The key advantage offered by this method is the highly polymorphic nature of the STR markers, which allow for a high probability of two-genome discrimination. Laboratories currently use different commercial multiplex kits for forensic identifications or in house assays, however have formed consortiums to standardize procedures and set guidelines for the correct interpretation of results, in order to improve this intrinsically semi-quantitative platform with a sensitivity of 1.0-3.0% (9).

Next generation sequencing (NGS) technologies are an innovation in human and animal genomics research, as they are capable of producing 100-fold more data than the most powerful Sanger based capillary sequencers; thus enabling researchers to investigate the large number of queries that remain to addressed (10).

NGS generates hundreds of giga-bases of nucleotide sequences per instrument run and produces this data at a lower cost, thus motivating researchers to use NGS for various purposes: To identify rare variations on the whole genome or on a target sequence, to analyze transcriptome profiling of cells, tissues and organisms and to identify epigenetic markers for disease diagnosis. Progress in the optimization of procedures, in addition to further reduction of costs, are the key factors that will lead to a more extensive uptake of this technique in diagnosis and for practical clinical applications.

NGS provides qualitative and quantitative data. Quantitative data depends on the depth of sequence data collected on each sample and on the quality of the target to expose. For samples with a lower abundance target, many more sequence reads are required to achieve accurate quantification (11). A previous study demonstrated that NGS exhibits sensitivity comparable to that of quantitative PCR (qPCR) in the evaluation of MRD in B cell disorders (12).

In the present study, an Ion AmpliSeq custom chimerism (ACCh) panel and a custom bioinformatic pipeline was created for chimerism quantification by NGS. The first aim was to detect the existence of cells of two origins in chimera samples and then to evaluate the capability of NGS to determine the percentage of the recipient cells.

Materials and methods

DNA sample preparation. The Ethics Committee of the Institute for Maternal and Child Health, IRCCS 'Burlo Garofolo' approved the present study (approval number: Prot. 18/2015, Cl. M/11). Written informed consent was obtained from all the participants.

Total peripheral blood was collected from 10 volunteer donors (V01-V10; 4 males and 6 females) with ages ranging from 20-50 years, and from 2 pediatric patients that underwent allogeneic HSCT (pR1, male, 5 years; and pR2, male, 12 years) and their donors (pD1, male, 9 years; and pD2, female, 25 years; Table I). Written informed consent was obtained from all the participants.

All DNA samples were isolated using the QIAamp DNA Blood kit according to manufacturer's protocol (Qiagen GmbH, Hilden, Germany). The DNA status was evaluated using the NanoDrop 1000 spectrophotometer (Thermo Fisher Scientific, Inc., Waltham, MA, USA) and the Qubit dsDNA HS Assay kit and the Qubit fluorometer (Thermo Fisher Scientific, Inc.). A DNA stock solution was prepared for all DNA samples at 20.0 ng/μ l.

DNA samples from the 10 volunteer donors were randomly paired. A total of 5 artificial chimeric DNA mixtures, as the donor/recipient chimera, were created by diluting DNA with its paired DNA at several percentages of the first DNA for each artificial DNA mixture (aCh).

Ion AmpliSeq custom chimerism panel design. A multi phase strategy was employed to evaluate the main characteristics of the ACCh panel: i) The panel average heterozygosity was assessed around 0.5 for the European population (HapMap Phase 3 CEU population); ii) two SNPs per somatic chromosome, termed 'main SNPs' (mSNPs), were selected and located in two different regions of the same chromosome; iii) the amplicon composition was evaluated according to the following requirements: a) GC percentage ranging between 40.0 and 60.0%; b) presence of one mSNP inside each amplicon; c) mSNP location preferably in the centre of the amplicon; d) absence of INDEL SNPs; e) absence of homopolymers and potential homopolymer generation from SNP variants and their flanking regions; f) absence of flanking SNPs to the mSNPs.

In total, 44 single-nucleotide, biallelic, polymorphisms were selected from the NCBI dbSNPs database (http://www.ncbi. nlm.nih.gov/SNP/, build 138, last database update 28.03.2014; Table II). A total of 4 base sequences including mSNPs were used as target regions for primer design. The primer pool, intended for DNA library construction through multiplex PCR, was defined by Ion AmpliSeq Designer software, version 3.0.1 (Thermo Fisher Scientific, Inc.). A single-tube, 44 primer pair pool was purchased from Life Technologies (Thermo Fisher Scientific, Inc.).

Ion torrent library preparation and sequencing. DNA sample library preparation was performed according to the AmpliSeq Library Preparation protocol (Life Technologies; Thermo Fisher Scientific, Inc.). For each DNA sample, a library was constructed using 10.0 ng genomic DNA through the Ion AmpliSeq Library kit, version 2.0. Subsequently, according to the library preparation protocol, each DNA library was indexed using the Ion Xpress Barcode Adapters kit (Thermo Fisher Scientific, Inc.) and was purified using AMPure XP magnetic beads (Beckman Coulter, Inc., Brea, CA, USA). Each DNA library was then quantified by qPCR using the thermo-cycler 7900HT Fast Real-Time PCR system with the Ion Library TQMN Quantification kit (Thermo Fisher Scientific, Inc.). Template Ion Sphere Particles were arranged using the Ion Personal Genome Machine (PGM) Template OT2 200 kit (Thermo Fisher Scientific, Inc.) and a single end 200 base-read sequencing run was conducted using the Ion Torrent PGM system. Libraries were pooled at 8 pM using the following rates: Donor/recipient, 1:1; and chimera/chimera, 1:1. The recipient/chimera rate was fixed at 1:40 in order to obtain an average coverage of the above libraries around 250X:10,000X. Library pools were sequenced on ion 314 and 316 chip (Table III).

Hotspot panel bed file. A hotspot panel bed file was created using the UCSC Genome Browser (https://genome.ucsc.edu). All SNPs located in the central region of each amplicon were included (NCBI dbSNPs build 138; 'Common SNPs' =286). All the INDELs present across the amplicons, and the SNPs near the 5' and 3' ends of the amplicons were excluded from the file. All the above 44 selected SNPs were marked as 'mSNP'

	DNA ID	Chimera ID	Chimera information	Notes
pD1	pR1	Samples used to evaluat	e panel informativity in consanguinity	Brothers, pre-HSCT
pD2	pR2	pCh ₁	MC by STR-CE analysis ^a	+1 month post-HSCT
pD2	pR2	pCh_2	CC by STR-CE analysis	+2 months post-HSCT
pD2	pR2	pCh ₃	MC by STR-CE analysis	+3 months post-HSCT
pD2	pR2	pCh_4	MC by STR-CE analysis	+4 months post-HSCT
pD2	pR2	pCh_5	MC by STR-CE analysis	+6 months post-HSCT
pD2	pR2	pCh_6	CC by STR-CE analysis	+10 months post-HSCT

Table I. List of the patient DNA samples used in the present study.

^aChimerism evaluation of all patient samples (pCh_{1-6}) was performed by STR-CE analysis in an external laboratory (Department of Molecular Medicine, University of Padova, Padova, Italy). HSCT, hematopoietic stem cell transplantation; MC, mixed chimerism; CC, complete chimerism; STR-CE, short tandem repeat capillary electrophoresis.

and the SNPs belonging to the same amplicon were indexed with the same chromosome/amplicon ID number. Finally, a hotspot panel bed file was created: 'HP286SNPs'.

Genotyping and quantification. Genotyping of all DNA was performed automatically, together with the quantification of all the chimeras, using a custom bioinformatics tool. The code of our tool was written using the Shiny package in R, a web framework to build interactive web applications (https://cran.r-project.org/web/packages/shiny/index.html). A functional diagram of the code is presented in Fig. 1 and the full code is available on request. The code is based on dependencies of the Bioconductor package that must be pre-installed for proper tool functionality.

The custom tool requires as input the sequencing bam files of 'Donor', ' Recipient' and 'Chimeric' patient. Briefly, it uses readGAlignments and pileLettersAt functions, from the GenomicAlignments package (13), to read bam files and extracts the letters/nucleotides into a set of individual genomic positions defined from the bed file. Thresholds for 'Donor' and 'Recipient', homozygous and heterozygous genotyping calls, are settled in base counts frequency ranges of 90.0-100.0% and 30.0-60.0%, respectively. Genotyping calls not included in the thresholds ranges were excluded as unreliable; users can modify the thresholds according to their needs from the user interface. Genotypes from each library were crosschecked to select only SNPs comparable in all conditions. Selected SNPs from donors and recipients were labelled as informative recipient alleles (IRA) according to the following schema: Donor homozygous and recipient heterozygous [Donor (AA) and Recipient (Aa); Donor (aa) and Recipient (Aa)]; donor and recipient homozygous for different alleles [Donor (AA) and Recipient (aa); Donor (aa) and Recipient (AA)].

Only the IRA SNPs tagged as informative were used to calculate the chimera's donor:recipient ratio as median of the allele frequency ratio, while standard error was used to calculate confidence intervals of prediction at 95.0%.

To cross validate the tool, genotyping of all donor and recipient samples was also performed manually obtaining the variant data from the Ion Torrent plugin Variant Caller, version 4.4 using the 'Generic-PGM-Germ Line-Low Stringency' configuration coupled by the HP286SNPs hotspot bed file.

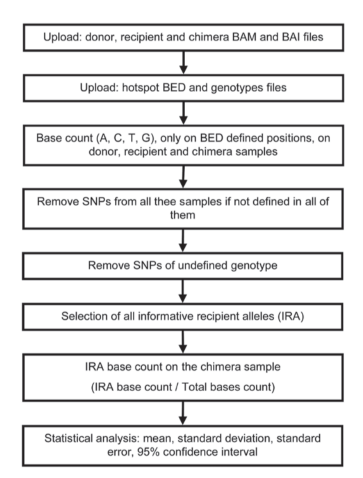


Figure 1. Code functional diagram. All output next generation sequencing files and reference files were uploaded to the custom pipeline. A quality control filtering step per target base was performed and the genotypes for donor and recipient samples were defined. Finally, subsequent to identification of the informative recipient allele, statistical analysis was conducted to determined the chimerism percentage. A, adenosine; C, cytosine; T, thymine; G, guanine; SNP, single nucleotide polymorphism; IRA, informative recipient allele.

Microsatellite analysis and patient data validation. Multiplex PCR amplification of V01-V10 and aCh₁₋₁₃ samples, in addition to the patient samples pD2, pR2 and pCh₁₋₆, was performed according to the manufacturer's instructions of the

Table II. List of all main SNPs included in the Ion AmpliSeq custom chimerism panel.

SNP ID	Genome position	Alleles	European heterozygosity	Informativity of recipient allele ^a %
rs12070036	chr1:g.227819514	A/G	0.407	41
rs1234315	chr1:g.173178463	C/T	0.513	37
rs10496711	chr2:g.134516742	C/G	0.407	40
rs12612347	chr2:g.219057338	A/G	0.442	40
rs1984630	chr3:g.134414219	G/T	0.522	36
rs9831477	chr3:g.30693522	A/T	0.483	38
rs10033900	chr4:g.110659067	C/T	0.496	37
rs5335	chr4:g.148463840	C/G	0.492	37
rs983889	chr5:g.15555486	A/C	0.487	38
rs10038113	chr5:g.25902342	C/T	0.469	38
rs552655	chr6:g.13370488	A/G	0.504	37
rs2077163	chr6:g.33636907	C/T	0.460	39
rs39395	chr7:g.103489729	A/G	0.425	40
rs2270188	chr7:g.116140524	G/T	0.496	38
rs10505477	chr8:g.128407443	C/T	0.531	36
rs532841	chr8:g.12957475	C/T	0.549	35
rs2297313	chr9:g.91669362	A/G	0.960	37
rs424539	chr9:g.14442595	C/G	0.467	38
rs1561570	chr10:g.13155726	C/T	0.522	36
rs619824	chr10:g.104581288	G/T	0.407	41
rs198464	chr11:g.61521621	C/T	0.504	37
rs178503	chr11:g.44082931	A/G	0.442	40
rs1126758	chr12:g.103248924	A/G	0.442	40
rs8608	chr12:g.53294381	A/G	0.522	36
rs1061472	chr13:g.52524488	A/G	0.522	37
rs504544	chr13:g.19735891	A/G A/T	0.508	37
rs10143250	chr14:g.104723433	C/T	0.434	39
rs1957779	chr14:g.63669647	C/T C/T	0.434	39
rs634990	chr15:g.35006073	A/G	0.449	39
rs2117215	chr15:g.94879684	C/T	0.603	32
rs121893	chr16:g.66183995	C/T C/T	0.414	32 39
rs2191125	chr16:g.7720923	T/C	0.550	34
rs6808	chr17:g.62400575	C/G	0.450	34
rs744166	chr17:g.40514201	T/C	0.430	40
	0	A/T		38
rs620898 rs633265	chr18:g.48509148		0.467	
	chr18:g.57831468	A/C	0.496	37
rs108295 rs892086	chr19:g.34224816	A/G C/T	0.496 0.451	37 39
	chr19:g.10837677			39 39
rs753381	chr20:g.39797465	T/C	0.451	
rs715147	chr20:g.50055350	G/A	0.367	43
rs225436	chr21:g.43729034	A/G	0.517	36
rs8128316	chr21:g.35721560	C/T	0.542	35
rs4444	chr22:g.31205334	A/G	0.483	38
rs132985	chr22:g.38563471	C/T	0.517	37

^aIn order to calculate the probability (informativity) to identify an informative recipient allele in a casual donor/recipient pair, the single values of probability of each genotype combination that carry an informative recipient allele were added as follows: Donor homozygous and recipient heterozygous [Donor (AA) and Recipient (Aa); Donor (aa) and Recipient (Aa)]; donor and recipient homozygous but for different alleles [Donor (AA) and Recipient (aa); Donor (aa) and Recipient (AA)]. SNP, single nucleotide polymorphism; A, adenine; G, guanine; C, cytosine; T, thymine.

DNA ID Chimera ID		Chimera notes	Ion chip
V01, V02	aCh_1	Chimera: 1.0% of V01; 99.0% of V02	314
V03, V04	aCh_2	Chimera: 1.25% of V03; 98.75% of V04	314
V03, V04	aCh ₃	Chimera: 2.5% of V03; 97.5% of V04	314
V05, V06	aCh_4	Chimera: 5.0% of V05; 95.0% of V06	314
V05, V06	aCh ₅	Chimera: 10.0% of V05; 90.0% of V06	314
V07, V08	aCh _{6,7,8,9,10,11}	Chimeras: 0.5%, 1.0%, 4.0%, 8.0%, 12.0%, 20.0% of V07; 99.5%, 99.0%, 96.0%, 92.0%, 88%, 80.0% of V08	316
V09, V10	aCh _{12,13}	Chimeras: 40.0% of V09 and 100% of V09; 60.0% of V10 and 0.0% of V10	316
pD2, pR2	pCh _{1,2,3,4,5,6}	Chimeras: MC, CC, MC, MC, MC, CC of pR2 ^a	316
pD2, pR2	pCh ₂	Technical replication	314
pD1, pR1	pCh ₆	Technical replication	314

Table III. List of the library pools sequenced by next generation sequencing.

^aAll patient samples were previously analyzed by short tandem repeat capillary electrophoresis in an external laboratory (Department of Molecular Medicine, University of Padova, Padova, Italy).

AmpFISTR Identifiler Plus PCR Amplification kit (Thermo Fisher Scientific, Inc.). Amplicons were resolved on a Genetic Analyzer 3130 and analyzed with GeneMapper software, version 4.1 (Life Technologies; Thermo Fisher Scientific, Inc.).

Patient samples (pCh₁₋₆) were also analyzed by qPCR (data not shown), as previously investigated by Bai *et al* (14). This analysis was performed as an additional validation method of NGS data, where a discrepancy between NGS and STR data was present.

Results

Ion chips and ACCh panel performance. A total of 7 library pools were loaded and sequenced on ion chip 314. Each pool was comprised of one donor, one recipient and one chimera. In addition, three additional pools were run on ion 316 chip (Table III). The mean values of performance of the 10 runs and of all samples are summarized in Table IV.

NGS genotyping performances using the ACCh panel. A total of 14 DNA samples were genotyped on Ion Torrent PGM using the ACCh panel with the HP286SNPs bed file. Regarding the mSNPs, the Variant Caller output identified that 2 mSNPs (rs121893 and rs12612347) were assigned as 'No Call' in over 50.0% of the genotyping runs due to low quality. The remaining 42 mSNPs were successfully genotyped. Concerning the remaining 242 SNPs, 27 SNPs were assigned as 'No Call', with an average of 10 SNPs per patient. The call of these SNPs failed in two Variant Caller filtering steps: 'Maximum common signal shift' and 'minimum coverage on either strand'.

Genotyping of all samples was additionally performed using our tool with the HP286SNPs bed file (Table V). To identify the IRAs, the data of the donor was compared with the recipient using the tool and manually cross-validated with the Variant Caller genotypes. Inside the genotyping calls of

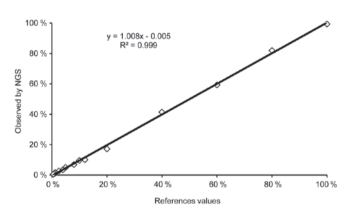


Figure 2. Linearity of Ion AmpliSeq custom chimerism panel: NGS against references values. A series of artificial chimeras, ranging between 0.5-100.0%, were quantified by NGS and plotted against their reference values. NGS, next generation sequencing.

242 SNPs, a small bias was present between these 2 tools; this is due to the high conserved filters of variant caller, dedicated predominantly for standard sequencing applications, and due to the absence of these filters in our custom tool.

NGS linearity, detection limit and accuracy with the ACCh panel. In order to test the linearity of Ion Torrent PGM with the ACCh panel in a fixed detection range (0.5-100.0%), a series of DNA mixtures was developed, diluting a DNA with its paired DNA at several percentages of the original. In order to increase the genetic marker variability in addition to the biological variability, a total of 12 artificial chimeras (aCh_{1-12}) were prepared from 5 different DNA pairs. Finally a pure DNA (V09) was run as 100.0% DNA (aCh_{13}). Subsequent to Ion Torrent sequencing, using the custom tool, quantitative data for all IRAs of each artificial chimera were obtained.

In addition, to increase the putative points in the dynamic range, the informative alleles of both DNA in the chimeras

A, In-house observed performance	Ion 314 chip	Ion 316 chip	
Ion sphere particles loading	84.3%	70.0%	
Total bases (Mb)	105.1	677.0	
Total reads	533,535	3,474,065	
Reads on-target	99.7%	99.3%	
Panel uniformity	97.6%	96.0%	
B, Coverage performance	Ion 314 chip	Ion 316 chip	
Chimera samples - amplicons over 2,500X	43/44	43/44	
	range 4,069X-23,944X)	(range 4,777X-28,314X)	
Donor & recipient samples - amplicons over 50X	44/44	44/44	
	(range 105X-952X)	(range 75X-2328X)	

Table IV. In-house Ion Torrent Personal Genome Machine analysis observed average performances using the Ion AmpliSeq custom chimerism panel on ion 314 and 316 chips.

	Custom	pipeline	IRA genotypes		
DNA pair 'Donor'/'Recipient'	44 mSNPs	242 SNPs	Heterozygous	Homozygous	Total IRAs (%) ^a
V01/V02	15	3	10	8	18 (43)
V03/V04	18	4	19	3	22 (52)
V05/V06	14	4	14	4	18 (43)
V07/V08	13	5	10	8	18 (43)
V09/V10	8	3	8	3	11 (26)
V02/V01	15	5	12	8	20 (48)
V04/V03	7	2	6	3	9 (21)
V06/V05	14	2	12	4	16 (38)
V08/V07	16	4	12	8	20 (48)
V10/V09	19	3	19	3	22 (52)
pD1/pR1	9	2	9	2	11 (26)
pD2/pR2	17	4	18	3	21 (50)

^aThe percentage of each IRA was calculated on 42-44 amplicons, due to the low quality of the remaining two, considering that each amplicon could contain at least one potential IRA. IRA, informative recipient allele; mSNP, main single nucleotide polymorphism.

 aCh_{12} (40.0% of V09 and 60.0% of V10) and aCh_{11} (20.0% of V07 and 80.0% of V08) were calculated and quantified.

Least-squares analysis of the above putative points identified a clear linearity ($R^2=0.999$; Y=1.008X-0.005) between NGS and the reference values (Fig. 2).

Analyzing the artificial chimeras aCh_{6-13} by capillary electrophoresis using the STRs markers, chimerism ranging from 4.0-100.0% was detected. Least-squares analysis identified a clear linearity (R²=0.999; Y=1.012X-0.009) between NGS and STRs values (Fig. 3).

In addition, the background of Ion Torrent generated by the ACCh panel was estimated. In this case, the custom tool was used, considering the 'donor' samples as chimera. The average background value at each SNP was estimated at 0.3% (range, 0.0-0.8%) and with a 95.0% confidence interval between 0.1 and 0.5%. Considering the background values and the reported literature on the error rates at each base of NGS technologies (range from 0.04-1.0%) (15), the detection limit of the NGS protocol with the ACCh panel was set at 1.0%, although an artificial chimera was detected at 0.5%.

Finally, considering that the method determined each chimera, calculating the average value of all IRA, the average standard error was used as an indirect marker of accuracy, using the data of all artificial chimeras ranging from 1.0-99.0% (excluding 0.5 and 100.0%). For the dynamic range of 1.0-20.0% of chimeras, the average standard error was

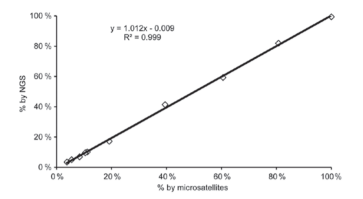


Figure 3. Linearity of Ion AmpliSeq custom chimerism panel: NGS against microsatellite values. A series of artificial chimeras were quantified by a standard microsatellite method (short tandem repeats) and the data were plotted compared to the NGS values. The chimeras ranging between 0.5-2.5% were not able to be quantified by short tandem repeats due to the detection limit of the informative markers. NGS, next generation sequencing.

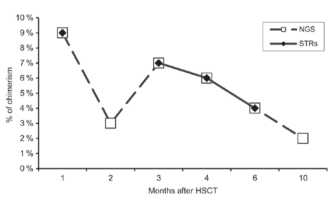


Figure 4. Patient sample evaluation with NGS and microsatellites (STRs) methods. Samples (n=6) taken from the same patient at different time points, ranging from 1-10 months subsequent to HSCT, were compared using the two methods. At +2 (pCh₂) and +10 (pCh₆) months, the STRs method was not able to detect the recipient DNA. NGS, next generation sequencing; STRs, short tandem repeats; HSCT, hematopoietic stem cell transplantation.

calculated at 0.3% with a deviation at 0.2%. For higher values of chimeras, up to 99.0%, the standard error increased up to 1.8% with a maximum deviation at 2.0%.

Patient chimerism evaluation on the NGS platform. Considering the linearity between NGS and reference values and between NGS and STR values using the standard NGS workflow and the custom tool, our workflow was tested in 6 samples of the same patient (pCh_{1-6}) in which the chimerism quantification report, previously performed by microsatellite analysis at different times in an external laboratory (Department of Molecular Medicine, University of Padova, Padova, Italy), presented at least one CC between two MCs (Table I).

NGS analysis detected a mixed chimerism in all samples $(pCh_1, pCh_2, pCh_3, pCh_4, pCh_5 and pCh_6)$ while the microsatellite analysis only in 4 of them $(pCh_1, pCh_3, pCh_4 and pCh_5)$. For these 4 MC samples, the percentage of predicted chimerism was equal between the 2 methods of analysis (Fig. 4). In regard to the pCh₂ and pCh₆ samples, NGS analysis evaluated a mixed chimerism at 3.0 and 2.0%, respectively (Fig. 4).

To confirm the obtained NGS data for the pCh₂ and pCh₆ samples, the NGS analysis was repeated, and qPCR was additionally perfomed in all patient samples. The results of qPCR analysis were in agreement with that of the NGS and microsatellite data for the pCh₁, pCh₃, pCh₄ and pCh₅, and confirmed the NGS data for the pCh₂ and pCh₆ samples.

Discussion

NGS technologies have revolutionized the field of genomics, and its application has been extended to different fields such as clinical diagnostics and forensic science (15-17). As a result of the continuous development of NGS, several applications previously performed on Sanger sequencing with capillary electrophoresis have been transferred onto the NGS platform, enabling fast and cost-effective generation sequence data with high resolution and accuracy. For this reason, different panels are being developed for the sequencing of genetic mutations involved in human diseases (e.g. MiSeqDx Cystic Fibrosis Clinical Sequencing Assay; Illumina, Inc., San Diego, CA, USA) or in cancer (Ion AmpliSeq BRCA1 and BRCA2 Panel; Life Technologies; Thermo Fisher Scientific, Inc.).

In the field of chimerism quantification by NGS platforms, Debeljak *et al* (18) reported an innovative and well performed study, using haplotype counting. In addition, Kim *et al* (19) briefly reported a relative quantification analysis of SNP markers by NGS in one human bone marrow chimerism sample. However, the study by Kim *et al* (19) was conducted in a 4.9% chimerism sample without any detail concerning the limit of detection, the technical error or additional important technical information and validation data of NGS application in chimerism quantification.

In the present study, a full workflow was designed, and the proposed protocols and a bioinformatics tool were tested for chimerism quantification by NGS. A 44-amplicon custom chimerism panel based on Ion AmpliSeq technology was designed, and in addition a bioinformatics tool dedicated to the genotyping and quantification of NGS data was coded. These resources were created in order to provide a novel tool for the evaluation of the chimerism following allogenic HSCT, thus potentially increasing the number of clinical analyses supported on NGS platforms.

The ACCh proposed panel is composed of 44 amplicons, containing 44 selected mSNPs, of which 2 mSNPs are located in different regions of each somatic chromosome. It is suggested that the different mSNP locations in all somatic chromosome may be useful to avoid predominantly false negatives results caused by chromosomal deletions characteristic of certain malignancies (20). In addition, the bed file uploaded in the custom tool, containing all targeted SNPs, can be modified in order to exclude the SNPs present in chromosome target regions subjected to deletions in a specific patient.

The panel average heterozygosity was assessed around 0.5 for the European population in order to obtain different informative markers for each transplanting pair, for a more precise and robust quantification. The theoretical panel informativity for unrelated donor:recipient values, calculated according to the data present on the NCBI dbSNPs database, was estimated to be approximately 16/42 mSNPs, while for siblings the informativity was estimated at 50.0% (approximately 8/42 mSNPs). In order to increase the informativity of the ACCh panel, an additional 242 selected SNPs present in the targeted regions were included in the bed file. This addition of SNPs experimentally increased the average informativity (Table V).

The ACCh panel reached the limit of detection on the Ion Torrent PGM platform of 0.5%, however, this was updated to the conservative value of 1.0% for two reasons: i) The Ion Torrent error is defined to be between 0.04 and 1.0% (21); and ii) the background of the ACCh panel, based on the IRA data of our experiments, ranges between 0.1 and 0.5%.

Regarding the timing of chimerism analysis, the UK NEQAS Consortium has recommended that results should be assessed in 5 working days from the reception of the sample and in 3 working days for urgent requests (6). The protocol suggested in the current study is feasible in 2 days; the first day for library preparation and quantification and the second for template preparation, run sequence and data analysis. In addition, due to the fact the ACCh protocol suggested in the current study does not present any differences from the standard AmpliSeq Library Preparation and Ion Torrent PGM Run Sequence protocols, it is possible to introduce it to a standard manual library preparation workflow or in a library preparation workstation.

Considering the AmpliSeq library construction protocol (based on multiplex PCR) from Life Technologies (Thermo Fisher Scientific, Inc.), the same concept could be tested on additional NGS platforms, such as Illumina or Roche, according to their library preparation protocols. Concerning the custom bioinformatics tool, any bam and bai file coupled by a bed file, generated from any platform could be used.

At present, the cost of NGS analysis, compared with microsatellite methods, remains high, however considering the continuously reducing cost per NGS run, an NGS-based method for chimerism quantification could be evaluated in the future for its adoption in laboratories with a high volume of activity, and with NGS platforms already in use for other purposes. Although the present study reported a clear correlation between NGS and STR methods and identifed important technical details, further experimental replications are required in order for the NGS protocol to be validated for future laboratory use.

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