

USE OF GENOTYPIC HIV DNA TESTING: A DELPHI-TYPE CONSENSUS Short running title: Delphi Consensus on Genotypic HIV DNA Testing

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2	A DELPHI-TYPE CONSENSUS
3	Short running title: Delphi Consensus on Genotypic HIV DNA Testing
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USE OF GENOTYPIC HIV DNA TESTING:

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1

30 Abstract

31 Objectives:

32 As many disparities in the clinical use of HIV DNA sequencing are observed, a DELPHI-type

33 consensus was initiated in France to homogenize use, techniques, and interpretation of results.

34 <u>Methods</u>:

Based on a literature review and clinical experience, a Steering Committee (SC) of eight
virologists and one infectious disease specialist formulated statements. Statements were
submitted to an independent and anonymous electronic vote of virologists and HIV clinicians
in France, between October and December 2022.

39 <u>Results</u>:

40 The SC developed 20 statements grouped into six categories: *clinical situations for the use of* 41 HIV DNA genotyping; techniques for performing HIV DNA genotyping; consideration of 42 APOBEC mutations; genotyping results reporting; recycling of antiretrovirals; availability of HIV DNA genotyping tests and delays. Twenty-one virologists and 47 clinicians participated 43 44 in two voting rounds and 18/20 (90%) assertions reached a 'strong' consensus. For example, that prior genotyping on HIV DNA is useful for clinical decision-making when considering 45 46 switching to some long-acting regimens or to reduce the number of antiretroviral agents in virologically suppressed patients for whom RNA data are unavailable / not exploitable / not 47 48 sufficiently informative. Two statements achieved no consensus: reporting any detected viral 49 minority population for discussion in multidisciplinary meetings (virologists), and possible risk of virologic failure when using a second generation InSTI + XTC regimen in patients 50 51 with undetectable viral load ≥ 1 year and in the presence of a documented *M184V* mutation <5 years (clinicians). 52

53 <u>Conclusion</u>:

54 This DELPHI-type consensus will facilitate the strengthening and harmonization of good 55 practice when performing HIV DNA sequencing.

56 **1. Introduction**

57 Human Immunodeficiency Virus-1 infection has become a manageable chronic disease with the availability of antiretroviral therapies.^{1, 2} Lifelong treatment is currently required to obtain 58 59 and maintain viral suppression. Either prior to initiation of ART or in the event of suboptimal response to ART, HIV drug resistance testing using plasma HIV RNA plays a key role in 60 guiding treatment choices and optimization.^{1,2} When switching to a new ART regimen due to 61 toxicities, for simplification, drug reduction, or a long-acting regimen, it is also recommended 62 to first check HIV genotyping data.² In these situations, HIV viral load (VL) usually under 50 63 copies/mL does not allow amplification for RNA drug resistance testing.³ 64

In recent years, there has been growing interest in how HIV drug resistance testing using 65 cellular HIV DNA could assist in clinical decision-making in the event of switching ART, 66 especially when plasma HIV RNA genotype testing is not possible.⁴⁻⁶ The 2022 European 67 68 AIDS Clinical Society (EACS) guidelines state that "Proviral DNA genotyping may be useful in persons with multiple virologic failures, unavailable resistance history or low-level viremia 69 at the time of switch".² European and French guidelines indicate that it is possible to perform 70 71 genotypic resistance tests on HIV DNA from Peripheral Blood Mononuclear Cells (PBMCs) in the absence of historical data on plasma viral RNA.^{2,7} This test should be interpreted with 72 caution since it has a good positive predictive value but a low negative predictive value.⁷ 73

However, while these guidelines provide general guidance on the indications for cellassociated total HIV DNA resistance testing, practical recommendations to virologists and
HIV clinicians are lacking, particularly regarding frequent specific ART switch situations,
technique, and interpretation of results.

Since many disparities in clinical practice have been observed, both in the literature and in clinical practice, a modified DELPHI consensus research project was conducted in France with the aim of homogenizing situations in which HIV DNA sequencing could be used andguiding interpretation of results.

82 **2.** Material and methods

The Delphi method is an iterative consensus approach based on information collected from a panel of voters with expertise in the subject under consideration.⁸⁻¹⁶ This approach has been widely used in many therapeutic areas and several times in HIV care.¹⁷⁻²⁸ Using this structured approach, voting experts give their opinion individually and anonymously, and express their degree of agreement on statements in order to achieve consensus on a specific and well-defined subject.

In accordance with both French and international methodologies,^{9-12,29} our study was structured as a modified national Delphi consensus and conducted among French hospital clinicians and virologists between September and December 2022. The opinion of voting experts was collected during two assessment rounds using a questionnaire developed by a Steering Committee (SC) (Figure 1).

As recommended by the French National Authority for Health (HAS), voters specified their level of agreement with the statements using a 9-point Likert scale ranging from 1 "Strongly disagree" to 9 "Strongly agree".²⁹⁻³¹ The percentage of scores and the median were calculated for each statement separately in each voting round. Consensus for a statement was considered 'strong' when >75% of the scores were \geq 7 and the median score was \geq 8, 'good' when only one of these two parameters was satisfied, and 'lacking' when none of the parameters was satisfied.^{9,10,32}

101 Steering Committee (SC)

102 The SC included one infectious disease specialist and eight virologists directed by the last103 author of this article. Two initial SC meetings were held in June and August 2022.

104 *Voting Group*

105 Two voter profiles were identified: virologists and HIV clinicians. A list of voters was 106 compiled based on the following criteria: experience, acquired knowledge and expertise in 107 HIV care, presenting in national conferences or involvement in HIV care projects, with 108 recruitment throughout France, including French overseas territories. The voters were invited 109 via individual e-mails to participate in online voting, with personalized access via a dedicated 110 website. Questions on techniques for performing HIV DNA genotyping were voted on by 111 virologists only. The anonymity of both voting groups was guaranteed. Voters had no interaction with the SC, and SC members did not vote.²⁹ 112

113 Voting Round #1

During this first round of voting, a free text space for comments was made available enabling voters to develop or explain their opinion for each statement. At the end of the first round, scores and voter comments were summarized for each statement.

117 A third SC meeting took place in November 2022 to discuss the Round #1 results:

Statements that achieved a 'strong' consensus (i.e., ≥75% of scores ≥7 AND median ≥8)
 were validated in full and included in the final summary.

- Statements that achieved a 'good' consensus (i.e., ≥75% of scores ≥7 OR median ≥8)
 were discussed and proposed for Voting Round #2 only when the SC was able to develop
 a revised version based on analysis of voter comments.
- Statements that did not achieve consensus were re-worded by the SC based on feedback
 from voters and submitted for Voting Round #2.

125 Voting Round #2

Only voters from Voting Round #1 were invited to participate in Voting Round #2 to assess the statements amended by the SC from Voting Round #1 results. The free text comment option was deleted but replaced with an 'I don't know' option instead of the scoring response. 129 Votes including this 'I don't know' option were excluded from the analysis. Following the130 results of Voting Round #2, the SC closed the process.

131 *Ethical considerations*

This research was conducted in accordance with the Declaration of Helsinki. All personal data
transmitted for the study was separated from the results and anonymized, pursuant to the
French data protection law (GDPR – General Data Protection Regulation).

135 **3. Results**

Based on a literature analysis, existing guidelines and clinical experience, the SC initially developed 21 statements (two were subsequently merged resulting in 20 statements) divided into 6 key areas: *clinical situations for the use of HIV DNA genotyping; techniques for performing HIV DNA genotyping; consideration of APOBEC mutations; genotyping results reporting; recycling of antiretrovirals; availability of HIV DNA genotyping tests and delays.*

141 *Participation*

142 Voters in Round #1 included 21 virologists and 47 clinicians. All virologists (21/21, 100%)
143 and 40 clinicians out of 47 (85.1%) from Round #1 actively voted in Round#2.

A summary of the characteristics of voters is shown in Table 1. The virologists were 76% (n=16) full-time hospital workers, 10% (n=2) part-time and 14% (n=3) engineers ('Others'). Their median experience in performing HIV DNA sequencing was 10 years (IQR [5-12]) and the median number of HIV DNA genotypes performed per year was 225 (IQR [42.5-425]). Clinicians were 94% (n=44) full time hospital workers and 6% (n=3) part time. Their median experience with people living with HIV (PLWH) management was 25 years (IQR [15-31.5]) and the median number of patients they followed per year was 270 (IQR [200-400]).

All virologists and clinicians had extensive experience in HIV care-related activities over the previous five years, such as writing conference abstracts (76% and 83% respectively), writing scientific publications (76% and 74%), participating in research projects (100% and 91%), involved in training (81% and 81%), belonging to a professional or associated group (76%
and 81%) and speaking at scientific events (52% and 68%).

156 Statements (Table 2)

157 After Voting Round #1, 9/21 statements achieved a 'strong' consensus (>75% votes >7 and 158 median ≥ 8); 5/21 statements achieved a 'good' consensus ($\geq 75\%$ votes ≥ 7 or median ≥ 8) and 159 7 statements lacked a consensus: 12 statements were revised by the SC for Voting Round #2, 160 including all those which achieved a 'good' consensus and all those which did not achieve a 161 consensus, of which two were merged resulting in 20 statements. After Voting Round #2, 9/11 revised statements achieved a 'strong' consensus, and two statements did not achieve a 162 consensus. In total, 18/20 statements (90%) achieved consensus. The distribution of 163 164 cumulative votes, medians and results are provided in Table 2.

165 Consensus statement results (See Table S1 for Consensus results according to voter group,
166 and Table S2 for Statements, detailed virologists and clinicians voting results, and cumulative
167 results for both groups).

168 <u>Clinical situations for the use of HIV DNA genotyping</u>

In the context of a therapeutic decision requiring genotyping data, there was a 'strong' consensus from voters on the recommendation to perform HIV DNA genotyping when HIV RNA is non-amplifiable, when cumulative HIV RNA genotyping is not available and/or when the historical genotype is incomplete or unusable. Voters recognized with a 'strong' consensus that for the following therapeutic targets - reverse transcriptase, protease, integrase - HIV DNA sequencing has a good positive predictive value towards mutation detection (excluding APOBEC mutations) and an imperfect negative predictive value.

176 Voters also 'strongly' agreed that, in a virologically suppressed patient, in the absence of
177 exploitable or sufficiently informative RNA data, and when considering a drug-reduction /
178 simplification of the antiretroviral (ARV) regimen:

- for a switch to some long-acting regimens, prior HIV DNA genotyping is useful for clinical
decision-making,

- for sequential dosing (4 days out of 7 or 5 days out of 7) without changing any ARV in the
current regimen, it is not mandatory to prior perform HIV DNA genotyping for clinical
decision-making,

- for a reduced ARV number regimen, prior HIV DNA genotyping may be useful for clinical
decision-making.

186 <u>Techniques for performing HIV-DNA genotyping</u>

Virologists validated with a 'strong' consensus that, in current practice, HIV DNA genotyping has a decreased performance (sensitivity, representativeness of viral populations) when the DNA quantity is very low. It can be performed indifferently from whole blood, mononuclear cells isolated from peripheral blood or blood cell pellets, and although performance could be increased by performing duplicate, duplicate is not feasible in clinical practice.

193 Concerning HIV DNA genotyping techniques, virologists agreed with a 'strong' consensus 194 that Sanger or ultra-deep sequencing (UDS) could be used. However, there was an absence of 195 consensus on the relevance of discussing any viral minority population (i.e. variants below 15 196 to 20% of the viral population) detected after using UDS techniques in a multidisciplinary 197 meeting in the absence of defined clinically relevant detection threshold, according to the 198 current state of knowledge ('no consensus'; with the exclusion of 1/21 (4.7%) virologists who 199 answered 'I don't know').

200

Consideration of APOBEC mutations

The cytidine deaminases APOBEC3F and 3G enzymes might introduce G to A nucleotide mutations that can impair crucial enzymatic sites or generate stop codons that reduce the amount of replication competent proviruses.³³⁻³⁶ Voters validated with a 'strong' consensus that the detection of the *M1841* mutation in HIV DNA is suggestive of the presence of a defective genome in the APOBEC enzyme when associated with other evocative mutations (e.g., *M411, M2301* on reverse transcriptase) and/or stop codons. They also recognized with a 'strong' consensus that, when resistance mutations attributable to APOBEC are present, their significance should be interpreted with caution according to the clinical context and therapeutic history of the patient and should be indicated in the HIV DNA genotyping analysis report.

211 <u>Reporting of genotyping results</u>

With a 'strong' consensus, virologists and clinicians felt that the clinical interpretation of resistance mutations on HIV DNA genotyping should be discussed in multidisciplinary meetings. They also agreed that the detection via HIV DNA sequencing of new resistance mutations (excluding APOBEC and stop codons), which were previously undetected, must be considered for the switch decision and subsequent patient follow-up.

217 <u>ARV recycling</u>

218 Virologists and clinicians agreed with a 'strong' consensus that, in a patient with an 219 undetectable viral load for at least one year and with documented M184V substitution on the 220 current DNA genotype and/or on an RNA genotype performed within the last five years, the use of a 2nd generation InSTI (Integrase Strand Transfer Inhibitor) + XTC (Lamivudine or 221 222 Emtricitabine) + 1 NRTI combination is at low risk of virological failure over time. 223 Virologists validated with a 'strong' consensus that, under the same conditions, the use of a 2^{nd} generation InSTI + XTC combination may present a risk of virological failure over time. 224 225 However, clinicians remained divided on this possible virological risk and their vote did not 226 reach a consensus ('no consensus', no clinicians answered 'I don't know').

With a 'strong' consensus, virologists and clinicians validated that the use/recycling of NNRTIs, if resistance to this class was detected in HIV-DNA and/or in previous historical genotypes, is associated with a greater risk of virological failure, independently of the duration of undetectable viral load, particularly in drug-reduction strategies using this ARVclass.

232

Availability of HIV DNA genotyping tests and time to report results

With a 'strong' consensus, virologists and clinicians felt that genotypic HIV DNA testing should be accessible in clinical practice to all clinicians managing PLWH, and that results from these tests should be available within 30 days.

4. Discussion

237 This consensus research, using the DELPHI method, aims at harmonizing HIV DNA238 sequencing practices.

239 All five assertions on clinical situations for use of HIV DNA genotyping developed by the SC 240 were validated with a 'strong' consensus by the voters. Although HIV DNA sequencing is not routinely recommended² and does not systematically reveal the same results as those 241 previously detected by cumulative plasma RNA genotyping in virologically controlled 242 243 patients,^{4,5} it is useful to perform in several clinical circumstances. This is the case when 244 historical HIV RNA resistance data are insufficient and/or incomplete, or when the viral load 245 is too low to proceed with HIV RNA sequencing. A recent study - based on a very large genotypic database in France - describing the prevalence of genotypic baseline risk factors for 246 some long-acting regimen failures among ARV-naive patients showed that 10.1% of patients 247 displayed one baseline virological risk factor for virologic failure.³⁷ These findings emphasize 248 249 the need to check the genotypic resistance profile prior to initiating a long-acting regimen to 250 limit the potential risk of virologic failure and the emergence of resistance.

However, in the case of a virologically suppressed patient, in the event of a decision to reduce or simplify sequential treatment (4 days or 5 days out of 7) without changing the regimen, there was a 'strong' consensus that prior genotyping of HIV DNA is not essential to clinical decision-making, even in the absence of usable or sufficiently informative RNA data. This 255 matches literature findings showing that triple combination therapy of a 2^{nd} generation InSTI 256 + XTC + 1 NRTI administered every 4 or 5 days maintains control of HIV replication in 257 virologically suppressed PLWH while reducing cumulative exposure to ARV.^{38,39}

258 There was a 'strong' consensus from virologist voters that HIV DNA sequencing should be 259 performed when the viral quantity is sufficiently high (since the quantity of HIV DNA 260 influences the quality of the results obtained), that it can be used from different blood sample matrix, indifferently by Sanger or UDS, and that duplicates increase test performance 261 262 (although this cannot be used in current clinical practice). Nevertheless, knowing HIV-DNA 263 genotyping underestimate resistance detection due to a phenomenon of dilution of resistant species in the reservoir regardless of the sequencing method used, UDS methods might 264 improve resistance detection in HIV-DNA due to their greater sensitivity.⁴⁰⁻⁴² Virologists 265 266 were unable to reach a consensus on the fact that, given the current state of knowledge, it may 267 be worthwhile reporting any minority viral population detected for discussion in a 268 multidisciplinary discussion. They also didn't support the idea that it might be useful to report 269 any minority viral population detected for multidisciplinary discussion in the current context 270 of an undefined detection threshold for UDS techniques.

271 Although the 1% threshold for UDS techniques was found to be close to the sensitivity obtained in historical HIV RNA resistance tests,⁴¹ it was difficult for the SC to generate a 272 statement for voting with such a detection threshold. This is due to the variability of this 273 274 threshold depending on the UDS technique used, and the lack of solid evidence on the impact 275 of a minority variant as low as 1% on virological failure for newer ARVs with a high barrier 276 to resistance. Considering that UDS on HIV-DNA is now affordable in clinical practice and 277 may become the future the potential new gold standard in the future, the definition of a 278 technical cut-off to warrant enough sequencing accuracy and a clinical cut-off to establish the 279 clinical relevance of minority variants on treatment switch in virologically suppressed patients 280 are still unmet needs. So further research into these thresholds for both RNA- and DNA-based 281 techniques is warranted.

As shown in the literature,^{43,44} the detection of the *M184I* mutation in HIV DNA suggests the 282 presence of a defective genome due to the APOBEC enzyme when associated with other 283 284 suggestive mutations (See Table S3 for the list of mutations)³⁶ and/or stop codons, and a 'strong' consensus was reached on this statement. The presence of M184I mutation can impair 285 286 the activity of XTC and possibly some nucleoside reverse transcriptase translocation 287 inhibitors (NRTTIs). These mutations should be considered possible artifacts if they occur at the same threshold at which multiple signature APOBEC mutations are also present.⁴⁵ When 288 289 resistance mutations attributable to APOBEC are detected, it is recommended that their significance should be interpreted with caution³⁷ and should be indicated in the HIV DNA 290 291 genotyping analysis.

The French National Authority for Health already recommends that the interpretation of results from a DNA-based genotypic resistance test requires consultation between clinician and virologist.⁴⁶ In this context, a 'strong' consensus was reached on the need to discuss clinical interpretation of resistance mutations obtained by HIV DNA genotyping at multidisciplinary discussions. This was also the case regarding clinical decisions about switching ART and patient follow-up in newly detected resistance mutations.

298 The question of how resistance mutations are 'archived' over time remains important for the 299 potential re-use of specific ARVs. A recent study investigated the kinetics of the M184V mutation in proviral HIV DNA in long-term virologically suppressed patients.⁴⁷ The authors 300 301 showed significant progressive clearance of the M184V mutation in proviral HIV DNA over 302 the five years of the study. In the presence of a detected M184V substitution over the past 5 303 years, the SC looked for consensus statements on ARV recycling practices. In this context, 304 the SC proposed statements on ARV recycling practices in the event of the presence of an 305 M184V substitution detected within the last five years.

Regardless of the finding of an *M184V* mutation in the DNA genotype and clearance kinetics of the mutation, it has been observed that, in patients virologically suppressed for at least one year, the use of a 2^{nd} generation InSTI + XTC + 1 NRTI regimen presents a low risk of 309 virological failure over time.⁴⁸ The voters 'strongly' endorsed this statement. However, when 310 a M184V mutation has been documented over the past five years in a virologically suppressed 311 patient, the virologist voters 'strongly' agreed that the use of a 2nd generation InSTI + XTC 312 regimen could present a risk of virological failure over time, as described in some literature.⁴⁹

For documented NNRTI mutations, there was a 'strong' consensus that the recycling of this ARV class is associated with an increased risk of virological failure, irrespective of the duration of viral suppression, particularly in drug reduction strategies and long-acting regimens using this ARV class.⁵⁰

Since HIV DNA sequencing adds an important contribution to many clinical situations and patient follow-up,² there was a 'strong' consensus that it should be accessible to all practitioners. Also, that its results should be received within one month. The literature rarely provides such an indication of time in which to report results but, with current HIV DNA sequencing methods being faster than before, this timeframe seems reasonable.⁵¹

322 The Delphi method is known as a structured procedure which enables many experts to be consulted individually and anonymously on a specific subject while guaranteeing free 323 324 expression of each voter. However, this approach has some limitations associated with voters' profiles, statements elaboration and criteria considered to achieve a consensus.⁵² Our research 325 326 sought to limit these potential biases as far as possible to ensure maximum objectivity. 327 Although voters were recruited only in France, they were selected on objective criteria based 328 on their experience and expertise in HIV care and HIV virology. These criteria yielded a voter 329 sample with reassuring characteristics: a median of 10 years' experience performing HIV 330 DNA sequencing in the virologists' group and a median of 25 years' experience in PLWH 331 management in the clinicians' group. As far as the SC statements are concerned, a literature 332 review made it possible to identify key questions raised in clinical practice and propose 333 precisely worded statements. In terms of the threshold used to reach consensus, our study was 334 based on a rigorous two-criteria approach. This strict and demanding definition lends a high 335 degree of credibility to our results. To ensure the virologist panel represented the whole of 336 France territory, the SC supported identification of some virologists. Finally, our research was conducted with a continuous and complete separation between voters who voted 337 338 anonymously and SC members who neither participated in the vote nor interacted directly with voters. The constraint inherent in this separation was the absence of direct exchanges 339 340 between voters and SC members: such exchanges could have been useful when revising 341 statements for voting Round #2. Furthermore, like all Delphi-type consensus, the findings 342 represent good practices for virologists and clinicians who remain masters of their own 343 practice and must adapt findings to individual patient circumstances.

In conclusion, in this consensus research using the Delphi method, 18/20 (90%) statements achieved a consensus. Only two assertions did not reach consensus. Virologist voters remained divided on the value of discussing any minority population detected at a multidisciplinary meeting, and the clinician voters remained divided on the possible virological risk of using a combination of a 2^{nd} generation InSTI + XTC in HIV suppressed patients of more than one year in the presence of a documented M184V mutation of less than five years.

Our consensus findings constitute a solid basis for implementation and homogenization of practice regarding the use of DNA HIV sequencing, its performance, and its reporting, particularly when needing to reduce the number of ARV agents and when using some longacting regimens.

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382 **References**

Panel on Antiretroviral Guidelines for Adults and Adolescents. Department of Health
 and Human Services. *Guidelines for the use of antiretroviral agents in adults and*

385		adolescents with HIV.
386		https://clinicalinfo.hiv.gov/sites/default/files/guidelines/documents/adult-adolescent-
387		arv/guidelines-adult-adolescent-arv.pdf
388	2.	European AIDS Clinical Society 2022. GUIDELINES Version 11.1.
389		https://www.eacsociety.org/media/guidelines-11.1_final_09-10.pdf
390	3.	Assoumou L, Charpentier C, Recordon-Pinson P et al. Prevalence of HIV-1 drug
391		resistance in treated patients with viral load .50 copies/mL: a 2014 French nationwide
392		study. J Antimicrob Chemother 2017; 72: 1769-1773.
393	4.	Wirden M, Soulie C, Valantin MA, et al. Historical HIV-RNA resistance test results
394		are more informative than proviral DNA genotyping in cases of suppressed or residual
395		viraemia. J Antimicrob Chemother 2011; 66: 709-12.
396	5.	Delaugerre C, Braun J, Charreau I, et al. Comparison of resistance mutation patterns
397		in historical plasma HIV RNA genotypes with those in current proviral HIV DNA
398		genotypes among extensively treated patients with suppressed replication. HIV Med
399		2012; 13 : 517-25.
400	6.	Boukli N, Boyd A, Collot M et al. Utility of HIV-1 DNA genotype in determining
401		antiretroviral resistance in patients with low or undetectable HIV RNA viral loads. J
402		Antimicrob Chemother 2018; 73: 3129-3136.
403	7.	Morlat, C. Groupe d'experts pour la prise en charge du VIH 2017. Optimisation d'un
404		traitement antirétroviral en situation de succès virologique. Prise en charge médicale
405		des personnes vivant avec le VIH. https://cns.sante.fr/actualites/prise-en-charge-du-
406		vih-recommandations-du-groupe-dexperts/
407	8.	Dalkey NC. The Delphi method: An experimental study of group opinion. In N. C.
408		Dalkey, D. L. Rourke, R. Lewis, & D. Snyder (Eds.). Studies in the quality of life:
409		Delphi and decision-making. Lexington, MA: Lexington Books, 1972; 13-54.

410	9.	Loblaw DA, Prestrud AA, Somerfield MR et al. American Society of Clinical
411		Oncology Clinical Practice Guidelines: formal systematic review-based consensus
412		methodology. J Clin Oncol 2012; 30 : 3136-3140.
413	10	. Boulkedid R, Abdoul H, Loustau M et al. Using and Reporting the Delphi Method for
414		Selecting Healthcare Quality Indicators: A Systematic Review. PLoS One 2011; 6:
415		e20476.
416	11	. Diamond IR, Grant RC, Feldman BM et al. Defining consensus: a systematic review
417		recommends methodologic criteria for reporting of Delphi studies. J Clin Epidemiol
418		2014; 67 : 401-9.
419	12	. Hasson F, Keeney S, McKenna H. Research guidelines for the Delphi survey
420		technique. J Adv Nurs 2000; 32: 1008-15.
421	13	. Hsu CC, Sandford BA. The Delphi Technique: Making Sense of Consensus. Practical
422		Assessment Research & Evaluation 2019; 12: 1-10.
423	14	. Humphrey-Murto S, Varpio L, Wood TJ et al. The Use of the Delphi and Other
424		Consensus Group Methods in Medical Education Research: A Review. Acad Med
425		2017; 92 : 1491-1498.
426	15	. Richard MA, Aubin F, Beneton N et al. Moderate Psoriasis in Clinical Practice:
427		French Expert Consensus Using a Modified Delphi Method. Adv Ther 2022; 39: 5203-
428		5215
429	16	. Kodjikian L, Baillif S, Couturier A et al. Recommendations for the management of
430		diabetic macular oedema with intravitreal dexamethasone implant: A national Delphi
431		consensus study. Eur J Ophthalmol 2022; 32: 2845-2856
432	17	. Tsui S, Denison JA, Kennedy CE et al. Identifying models of HIV care and treatment
433		service delivery in Tanzania, Uganda, and Zambia using cluster analysis and Delphi
434		survey. BMC Health Serv Res 2017; 17: 811.

435	18. Johnson MO, Neilands TB, Koester KA, et al. Detecting Disengagement From HIV
436	Care Before It Is Too Late: Development and Preliminary Validation of a Novel Index
437	of Engagement in HIV Care. J Acquir Immune Defic Syndr 2019; 81: 145-152.
438	19. O'Connell KA, Kisteneff AV, Gill SS et al. HIV post-exposure prophylaxis in the
439	emergency department: An updated assessment and opportunities for HIV prevention
440	identified. Am J Emerg Med 2021; 46: 323-328.
441	20. Fredericksen RJ, Edwards TC, Merlin JS et al. Patient and provider priorities for self-
442	reported domains of HIV clinical care. AIDS Care 2015; 27: 1255-64
443	21. Cummins D, Waters D, Aggar C et al. Assessing Risk of HIV-Associated
444	Neurocognitive Disorder. Nurs Res 2019; 68: 22-28.
445	22. Greacen T, Kersaudy-Rahib D, Le Gall JM et al. Comparing the Information and
446	Support Needs of Different Population Groups in Preparation for 2015 Government
447	Approval for HIV Self-testing in France. PLoS One 2016; 11: e0152567.
448	23. Feyissa GT, Lockwood C, Woldie M et al. Evaluation of a guideline developed to
449	reduce HIV-related stigma and discrimination in healthcare settings and establishing
450	consensus. <i>PLoS One</i> 2018; 13 : e0198781.
451	24. Johnson MO Koester KA, Wood T et al. Development of an Index of Engagement in
452	HIV Care: An Adapted Internet-Based Delphi Process. JMIR Res Protoc 2017; 6:
453	e224.
454	25. Uyei J, Li L, Braithwaite RS. HIV and Alcohol Research Priorities of City, State, and
455	Federal Policymakers: Results of a Delphi Study. Am J Public Health 2015; 105: e23-
456	6.
457	26. Maserati R, Antinori A, Bonora S et al. Optimizing HIV therapy. A consensus project
458	on differences between cytidine analogues and regime compactness. New Microbiol
459	2014; 37 : 285-306
460	27. Adegbehingbe SM, Paul-Ebhohimhem V, Marais D. Development of an AFASS
461	assessment and screening tool towards the prevention of mother-to-child HIV

462	transmission (PMTCT) in sub-Saharan Africa-a Delphi survey. BMC Public Health
463	2012; 12 : 402.
464	28. Engler K, Ahmed S, Lessard D et al. Assessing the Content Validity of a New Patient-
465	Reported Measure of Barriers to Antiretroviral Therapy Adherence for Electronic
466	Administration in Routine HIV Care: Proposal for a Web-Based Delphi Study. JMIR
467	<i>Res Protoc</i> 2019; 8 : e12836.
468	29. HAS (Haute Autorité de Santé) Guide méthodologique. 2010. Élaboration de
469	recommandations de bonne pratique. Méthode « Recommandations pour la pratique
470	clinique ». https://www.has-sante.fr/upload/docs/application/pdf/2011-
471	01/guide_methodologique_consensus_formalise.pdf
472	30. Letrilliart L, Milliat-Guittard L, Romestaing P et al. Building a shared patient record
473	for breast cancer management: a French Delphi study. Eur J Cancer Care (Engl)
474	2009; 18 : 131-9.
475	31. McMillan SS, King M, Tully MP. How to use the nominal group and Delphi
476	techniques. Int J Clin Pharm 2016; 38: 655-62.
477	32. Koene S, van Bon L, Bertini E et al. Outcome measures for children with
478	mitochondrial disease: consensus recommendations for future studies from a Delphi-
479	based international workshop. J Inherit Metab Dis 2018; 41: 1267-1273
480	33. Mangeat B, Turelli P, Caron G et al. Broad antiretroviral defence by human
481	APOBEC3G through lethal editing of nascent reverse transcripts. Nature 2003; 424:
482	99–103
483	34. Armitage AE, Deforche K, Chang CH et al. APOBEC3G-induced hypermutation of
484	human immunodeficiency virus type-1 is typically a discrete "all or nothing"
485	phenomenon. PLos Genet 2012; 8: e1002550.
486	35. Russell RA, Moore MD, Hu WS et al. APOBEC3G induces a hypermutation gradient:
487	purifying selection at multiple steps during HIV-1 replication results in levels of G-to-

488		A mutations that are high in DNA, intermediate in cellular viral RNA, and low in
489		virion RNA. <i>Retrovirology</i> 2009; 6: 16.
490	36.	Armenia D, Gagliardini R, Alteri et al. Temporal trend of drug-resistance and
491		APOBEC editing in PBMC genotypic resistance tests from HIV-1 infected
492		virologically suppressed individuals. J Clin Virol 2023; 168: 105551.
493	37.	Charpentier C, Storto A, Soulié C et al. Prevalence of genotypic baseline risk factors
494		for cabotegravir + rilpivirine failure among ARV-naive patients. J Antimicrob
495		Chemother 2021; 76 : 2983-2987.
496	38.	Sellem B, Abdi B, Lê M et al. Intermittent Bictegravir/Emtricitabine/Tenofovir
497		Alafenamide Treatment Maintains High Level of Viral Suppression in Virally
498		Suppressed People Living with HIV. J Pers Med 2023; 13: 583.
499	39.	Landman R, de Truchis P, Assoumou L, et al. A 4-days-on and 3-days-off
500		maintenance treatment strategy for adults with HIV-1 (ANRS 170 QUATUOR): a
501		randomised, open-label, multicentre, parallel, non-inferiority trial. Lancet HIV 2022;
502		9 : e79-e90.
503	40.	St John EP, Simen BB, Turenchalk GS et al. A Follow-Up of the Multicenter
504		Collaborative Study on HIV-1 Drug Resistance and Tropism Testing Using 454 Ultra
505		Deep Pyrosequencing. <i>PLoS ONE</i> 2016; 11 : e0146687.
506	41.	Rodriguez C, Nere ML, Demontant V et al. Ultra-deep sequencing improves the
507		detection of drug resistance in cellular DNA from HIV-infected patients on ART with
508		suppressed viraemia. J Antimicrob Chemother 2018; 73: 3122-3128.
509	42.	Balakrishna S, Loosli T, Zaheri M et al. Frequency matters: comparison of drug
510		resistance mutation detection by Sanger and next-generation sequencing in HIV-1. J
511		Antimicrob Chemother 2023; 78: 656–664.
512	43.	Charpentier C., Montes B., Perrier M et al. HIV-1 DNA ultra-deep sequencing
513		analysis at initiation of the dual therapy dolutegravir + lamivudine in the maintenance
514		DOLULAM pilot study. J Antimicrob Chemother 2017; 72: 2831-2836.

515 44. Allavena C, Rodallec A, Leplat A, et al. Interest of proviral HIV-1 DNA genotypic 516 resistance testing in virologically suppressed patients candidate for maintenance 517 therapy. J Virol Methods 2018; 251: 106-110. 518 45. Tzou PL, Kosakovsky Pond SL, Avila-Rios S et al. Analysis of unusual and signature 519 APOBEC-mutations in HIV-1 pol next-generation sequences. PLoS One 2020; 15: 520 e0225352. 521 46. Morlat, C. Groupe d'experts pour la prise en charge du VIH. Résistance du VIH-1 aux 522 antirétroviraux. 2016. Prise en charge médicale des personnes vivant avec le VIH. 523 https://cns.sante.fr/wp-content/uploads/2017/02/experts-vih_resistance.pdf 524 47. Palich R, Teyssou E, Sayon S et al. Kinetics of Archived M184V Mutation in 525 Treatment-Experienced Virally Suppressed HIV-Infected Patients. J Infect Dis 2022; 526 225: 502-509. 527 48. Andreatta K, Willkom M, Martin R et al. Switching to 528 bictegravir/emtricitabine/tenofovir alafenamide maintained HIV-1 RNA suppression 529 in participants with archived antiretroviral resistance including M184V/I. J 530 Antimicrob Chemother 2019; 74: 3555-3564. 531 49. Santoro MM, Armenia D, Teyssou E et al. Virological efficacy of switch to DTG plus 532 3TC in a retrospective observational cohort of suppressed HIV-1 patients with or 533 without past M184V: the LAMRES study. J Glob Antimicrob Resist 2022; 31: 52-62. 534 50. Marcelin AG, Soulie C, Wirden M et al. The Virostar study: analysis of emergent 535 resistance-associated mutations at first- or second-line HIV-1 virological failure with 536 second-generation InSTIs in two- and three-drug regimens. HIV Glasgow congress 537 2022. Abstract P225. 538 51. Alidjinou EK, Deldalle J, Hallaert C et al. RNA and DNA Sanger sequencing versus 539 next-generation sequencing for HIV-1 drug resistance testing in treatment-naive 540 patients. Antimicrob Chemother 2017; 72: 2823-2830.

- 541 52. Skinner R, Nelson RR, Chin WW *et al.* The Delphi Method Research Strategy in
- 542 Studies of Information Systems. *Commun Assoc Inf Syst* 2015; **37**: 31-63.

Figure 1: Modified Delphi process chart



AND median score = 8 or 9

 $OR median \ score = 8 \ or 9$



Characteristic	Virologists $(n = 21)$	Clinicians (n = 47)		
Age, median [IQR], years	46 [43-55]	56 [46-60.5]		
Gender F/M, n (%)				
Female	16 (76)	18 (38)		
Male	5 (24)	29 (62)		
Type of practice, n (%)				
Full-time hospital workers	16 (76)	44 (94)		
Part-time hospital workers	2 (10)	3 (6)		
Others	3 (14)	-		
Years of experience performing HIV DNA sequencing, median [IQR], years	10 [5-12]	-		
Number of HIV DNA genotypes performed per year, median [IQR]	225 [42.5-425]	-		
Years of experience in PLWH management, median [IQR], years	-	25 [15-31.5]		
Number of PLWH seen per year, median [IQR]	-	270 [200-400]		
Experience in HIV care-related activities in the past 5 years, n (%)				
Conference abstract	16 (76)	39 (83)		
Scientific article	16 (76)	35 (74)		
Research project (not including this study)	21 (100)	43 (91)		
Involved in training	17 (81)	38 (81)		
Professional or associate group or member	16 (76)	38 (81)		
Speaker at scientific events	11 (52)	32 (68)		

DNA: desoxyribonucleic acid; F: Female; M: Male; PLWH: People living with HIV

Table 2 - Statements and cumulative voting results for virologists and clinicians

For each statement, a total number of voters equaling 21 indicates that only virologists were invited to vote and a total number of voters different from 68, 61, or 21 indicates the use of the 'I don't know' option by voters during the second voting round.

	STATEMENTS	Scores 1-2-3 (n)	Scores 4-5-6 (n)	Scores 7-8-9 (n)	Median	Results
Clini	cal situations for the use of HIV DNA genotyping					
1	In the context of a therapeutic decision requiring genotyping data, when HIV RNA is not amplifiable, when cumulative HIV RNA genotyping is not available and/or or in the event of an incomplete or unusable genotypic history, HIV DNA genotyping is recommended.	2.9% (2)	7.4% (5)	89.7% (61)	9	Strong consensus
2	For the following therapeutic targets - reverse transcriptase, protease, integrase - HIV DNA sequencing has a good positive predictive value (excluding APOBEC mutations) and an imperfect negative predictive value.	2.9% (2)	13.2% (9)	83.8% (57)	8	Strong consensus
3	In the context of a patient who has achieved virological success, in the event of a decision to reduce or simplify treatment to some long-acting regimens, prior genotyping on HIV DNA is useful for clinical decision-making in the absence of usable or sufficiently informative RNA data.	5.9% (4)	13.2% (9)	80.9% (55)	8	Strong consensus
4	In the context of a patient with virological success, in the event of a decision to reduce or simplify sequential treatment (4 days out of 7 or 5 days out of 7) without changing the treatment molecules, prior genotyping on HIV DNA is not essential for the clinical decision, even in the absence of usable or sufficiently informative RNA data.	13.3% (8)	13.3% 10% (8) (6)		8	Strong consensus
5	In the absence of usable or sufficiently informative RNA data, in the case of a patient with virological success, in the event of a decision to reduce/simplify to a treatment that reduces the number of ARV, prior genotyping on HIV DNA may be useful for clinical decision-making.	8.2% (5)	13.1% (8)	78.7% (48)	8	Strong consensus
Tech	niques for performing HIV DNA genotyping					
6	In current practice, when the amount of DNA is very low, the performance (sensitivity, representativeness of viral populations) of HIV DNA genotyping is reduced.	5.3% (1)	5.3% (1)	89.5% (17)	9	Strong consensus
7	In current practice, genotyping on HIV DNA can be performed either from whole blood, from the cells isolated from peripheral blood (PBMC) or from blood cell pellets.	14.3% (3)	9.5% (2)	76.2% (16)	8	Strong consensus
8	Duplicate DNA genotyping increases performance (sensitivity, representativeness of viral populations) but is not possible in current practice.	10.5% (2)	5.3% (1)	84.2% (16)	8	Strong consensus
9	Sanger and ultra-deep sequencing can be used to perform HIV DNA genotyping.	0% (0)	23.8% (5)	76.2% (16)	8	Strong consensus
10	For ultra-high throughput DNA sequencing techniques (ultra- deep sequencing), with the current state of knowledge, the clinically relevant detection threshold is not defined. Nevertheless, it may be interesting to report any viral minority population detected for multidisciplinary discussions.	10% (2)	25% (5)	65% (13)	7.5	NO CONSENSUS
Cons	ideration of APOBEC mutations					
11	Detection of the <i>M1841</i> mutation in HIV DNA is suggestive of the presence of a defective genome due to the APOBEC enzyme when it is associated with other suggestive mutations (e.g., <i>M411</i> , <i>M2301</i> on reverse transcriptase) and/or stop codons.	1.9% (1)	13.2% (7)	84.9% (45)	8	Strong consensus
12	When resistance mutations attributable to APOBEC are present, their significance should be interpreted with caution and based on the clinical context and treatment history.	5.2% (3)	5.2% (3)	89.7% (52)	9	Strong consensus
13	The presence of resistance mutations attributable to APOBEC should be reported in the HIV DNA genotyping analysis.	9.5% (2)	4.8% (1)	85.7% (18)	8	Strong consensus
Repo	rting of genotyping results					
14	The clinical interpretation of resistance mutations on HIV DNA genotyping should be discussed at multidisciplinary consultation meetings.	4.4% (3)	17.2% (10)	80.9% (55)	8	Strong consensus
15	In a patient with virological success, the detection on HIV DNA genotyping of new resistance mutations (excluding APOBEC & stop codons) previously undetected must be considered for the	5% (3)	10% (6)	85% (51)	8	Strong consensus

	switch decision and subsequent follow-up.									
ARV	ARV recycling									
16	In a patient with an undetectable viral load for at least one year, in the presence of a documented M184V substitution over the past five years, the use of a 2^{nd} generation InSTI + XTC + 1 NRTI combination presents low risk of virological failure over time.	7.4% (5)	7.4% (5)	85.3% (58)	8	Strong consensus				
17	In a patient with an undetectable viral load for at least one year, in the presence of a documented M184V substitution over the past five years, the use of a 2^{nd} generation InSTI + XTC combination might present a risk of virological failure over time.	21.3% (13)	14.8% (9)	63.9% (39)	7	NO CONSENSUS				
18	The use / recycling of NNRTIs in the event of documented resistance to ARV of this class is associated with a greater risk of virological failure, independently of the duration of undetectable viral load, especially in drug-reduction strategies using this ARV class.	1.7% (1)	18.3% (11)	80% (48)	8	Strong consensus				
Test	Test availability and delays									
19	HIV DNA genotypic tests should be accessible in clinical practice to all clinicians managing PLWH.	8.8% (6)	7.4% (5)	83.8% (57)	9	Strong consensus				
20	Reports of genotypic HIV DNA test results should be sent to clinicians within 30 days.	6.6% (4)	9.8% (6)	83.6% (51)	9	Strong consensus				

ARV: Antiretrovirals; APOBEC: Apolipoproteins B mRNA editing enzyme; InSTI: Integrase Strand Transfer Inhibitor; XTC: Lamivudine or Emtricitabine; PLWH: People living with HIV.

Table S1 – Consensus results according to voter groups

	STATEMENTS	Results VIROLOGISTS	Results CLINICIANS	CUMULATIVE Results
Clini	cal situations for the use of HIV DNA genotyping			
1	In the context of a therapeutic decision requiring genotyping data, when HIV RNA is not amplifiable, when cumulative HIV RNA genotyping is not available and/or or in the event of an incomplete or unusable genotypic history, HIV DNA genotyping is recommended.	Strong consensus	Strong consensus	STRONG CONSENSUS
2	For the following therapeutic targets - reverse transcriptase, protease, integrase - HIV DNA sequencing has a good positive predictive value (excluding APOBEC mutations) and an imperfect negative predictive value.	Strong consensus	Strong consensus	STRONG CONSENSUS
3	In the context of a patient who has achieved virological success, in the event of a decision to reduce or simplify treatment to some long-acting regimens, prior genotyping on HIV DNA is useful for the clinical decision in the absence of usable or sufficiently informative RNA data.	Strong consensus	Strong consensus	STRONG CONSENSUS
4	In a virologically suppressed patient, in the event of a decision to reduce or simplify to a sequential treatment (4 days out of 7 or 5 days out of 7) without changing any ARV agents, prior genotyping on HIV DNA is not essential for clinical decision- making, even in the absence of usable or sufficiently informative RNA data.	Strong consensus	Good consensus	STRONG CONSENSUS
5	In the absence of usable or sufficiently informative RNA data in a virologically suppressed patient, in the event of a decision to reduce/simplify to a treatment that reduces the number of ARV, prior genotyping on HIV DNA may be useful for clinical decision-making.	Strong consensus	Strong consensus	STRONG CONSENSUS
Tech	niques for performing HIV DNA genotyping			
6	In current practice, when the amount of DNA is very low, the performance (sensitivity, representativeness of viral populations) of HIV DNA genotyping is reduced.	Strong consensus	N.A.	STRONG CONSENSUS
7	In current practice, genotyping on HIV DNA can be performed either from whole blood, from the cells isolated from peripheral blood (PBMC) or blood cell pellets.	Strong consensus	N.A.	STRONG CONSENSUS
8	Duplicate DNA genotyping increases performance (sensitivity, representativeness of viral populations) but is not possible in current clinical practice.	Strong consensus	N.A.	STRONG CONSENSUS
9	Sanger and ultra-deep sequencing can be used to perform HIV DNA genotyping.	Strong consensus	N.A.	STRONG CONSENSUS
10	For ultra-high throughput DNA sequencing techniques (ultra- deep sequencing), with the current state of knowledge, the clinically relevant detection threshold is not defined. Nevertheless, it may be interesting to report any viral minority population detected for multidisciplinary discussions.	No Consensus	N.A.	NO CONSENSUS
Cons	ideration of APOBEC mutations			
11	Detection of the <i>M1841</i> mutation in HIV DNA is suggestive of the presence of a defective genome due to the APOBEC enzyme when it is associated with other suggestive mutations (e.g., <i>M411</i> , <i>M2301</i> on reverse transcriptase) and/or stop codons.	Strong consensus	Strong consensus	STRONG CONSENSUS
12	When resistance mutations attributable to APOBEC are detected, their significance should be interpreted with caution and based on the clinical context and treatment history.	Strong consensus	Strong consensus	STRONG CONSENSUS
13	The presence of resistance mutations attributable to APOBEC should be reported in the HIV DNA genotyping analysis.	Strong consensus	N.A.	STRONG CONSENSUS
Repo	rting of genotyping results			
14	The clinical interpretation of resistance mutations on HIV DNA genotyping should be discussed at multidisciplinary consultation meetings.	Strong consensus	Strong consensus	STRONG CONSENSUS
15	In a virologically suppressed patient, the detection on HIV DNA genotyping of new resistance mutations (excluding APOBEC & stop codons) previously undetected must be	Strong consensus	Strong consensus	Strong consensus

	considered for the treatment switch decision and subsequent									
	follow-up.									
ARV	ARV recycling									
16	In a patient with an undetectable viral load for at least one year, in the presence of a documented M184V substitution over the past five years, the use of a 2^{nd} generation InSTI + XTC + 1 NRTI combination presents low risk of virological failure over time.	Strong consensus	Strong consensus	STRONG CONSENSUS						
17	In a patient with an undetectable viral load for at least one year, in the presence of a documented M184V substitution over the past five years, the use of a 2^{nd} generation InSTI + XTC combination might present a risk of virological failure over time.	Strong consensus	No consensus	NO CONSENSUS						
18	The use / recycling of NNRTIs in the event of documented resistance to ARV agents of this class is associated with a greater risk of virological failure, independently of the duration of undetectable viral load, especially in drug-reduction strategies using ARV agents of this class.	Strong consensus	Strong consensus	STRONG CONSENSUS						
Test	Test availability and delays									
19	HIV DNA genotypic tests should be accessible in clinical practice to all clinicians managing PLWH.	Strong consensus	Strong consensus	STRONG CONSENSUS						
20	Reports of genotypic HIV DNA test results should be given to clinicians within 30 days.	Strong consensus	Strong consensus	STRONG CONSENSUS						

N.A.: Not available; ARV: Antiretroviral; APOBEC: Apolipoproteins B mRNA editing enzyme; InSTI: Integrase Strand Transfer Inhibitor; XTC: Lamivudine/Emtricitabine; PLWH: People living with HIV.

Table S2 – Statements, detailed virologists' and clinicians' voting results, and cumulative results for both groups

For each statement, a total number of voters different from 21 for virologists and 40 or 47 for clinicians indicates the use of the 'I don't know' option by voters during the second voting round.

		VOTING SCORES FROM VIROLOGISTS				VOTING SCORES FROM CLINICIANS					CUMULATIVE	
	STATEMENTS	Scores 1-2-3 (n)	Scores 4-5-6 (n)	Scores 7-8-9 (n)	Median	Results	Scores 1-2-3 (n)	Scores 4-5-6 (n)	Scores 7-8-9 (n)	Median	Results	SCORES
Clin	nical situations for the use of HIV DNA genotyping	5										
1	In the context of a therapeutic decision requiring genotyping data, when HIV RNA is not amplifiable, when cumulative HIV RNA genotyping is not available and/or or in the event of an incomplete or unusable genotypic history, HIV DNA genotyping is recommended.	4.8% (1)	0% (0)	95.2% (20)	9	Strong consensus	2.1% (1)	10.6% (5)	87.2% (41)	8	Strong consensus	STRONG CONSENSUS
2	For the following therapeutic targets - reverse transcriptase, protease, integrase - HIV DNA sequencing has a good positive predictive value (excluding APOBEC mutations) and an imperfect negative predictive value.	4.8% (1)	4.8% (1)	90.5% (19)	9	Strong consensus	2.1% (1)	17.0% (8)	80.9% (38)	8	Strong consensus	STRONG CONSENSUS
3	In the context of a patient with virological success, in the event of a decision to reduce or simplify treatment to some long-acting regimens, prior genotyping on HIV DNA is useful for clinical decision-making in the absence of usable or sufficiently informative RNA data.	9.5% (2)	4.8% (1)	85.7% (18)	9	Strong consensus	6.4% (3)	14.9% (7)	78.7% (37)	8	Strong consensus	STRONG CONSENSUS
4	In a virologically suppressed patient, in the event of a decision to reduce or simplify to a sequential treatment (4 days out of 7 or 5 days out of 7) without changing any ARV agents, prior genotyping on HIV DNA is not essential for the clinical decision, even in the absence of usable or sufficiently informative RNA data.	10% (2)	5% (1)	85.0% (17)	8	Strong consensus	15% (6)	12.5% (5)	72.5% (29)	8	Good consensus	STRONG CONSENSUS
5	In the absence of usable or sufficiently informative RNA data in a virologically suppressed patient, in the event of a desire to reduce/simplify towards a regimen that reduces the number of ARV agents prior genotyping on HIV DNA may be useful for clinical decision-making.	9.5% (2)	4.8% (1)	85.7% (18)	8	Strong consensus	7.5% (3)	17.5% (7)	75.0% (30)	8	Strong consensus	STRONG CONSENSUS
Tec	hniques for performing HIV DNA genotyping											
6	In current practice, when the amount of DNA is very low, the performance (sensitivity, representativeness of viral populations) of HIV DNA genotyping is reduced.	5.3% (1)	5.3% (1)	89.5% (17)	9	Strong consensus	N.A.	N.A.	N.A.	N.A.	N.A.	STRONG CONSENSUS
7	In current practice, genotyping on HIV DNA can be performed either from whole blood, from the cells isolated from peripheral blood (PBMC) or from blood	14.3% (3)	9.5% (2)	76.2% (16)	8	Strong consensus	N.A.	N.A.	N.A.	N.A.	N.A.	STRONG CONSENSUS

	cell pellets.											
8	Duplicate DNA genotyping increases performance (sensitivity, representativeness of viral populations) but is not possible in current clinical practice.	10.5% (2)	5.3% (1)	84.2% (16)	8	Strong consensus	N.A.	N.A.	N.A.	N.A.	N.A.	STRONG CONSENSUS
9	Sanger and ultra-deep sequencing can be used to perform HIV DNA genotyping.	0% (0)	23.8% (5)	76.2% (16)	8	Strong consensus	N.A.	N.A.	N.A.	N.A.	N.A.	STRONG CONSENSUS
10	For ultra-high throughput DNA sequencing techniques (ultra-deep sequencing), with the current state of knowledge, the clinically relevant detection threshold is not defined. Nevertheless, it may be interesting to report to report any viral minority population detected for multidisciplinary discussions.	10% (2)	25% (5)	65% (13)	7.5	No Consensus	N.A.	N.A.	N.A.	N.A.	N.A.	
Con	sideration of APOBEC mutations											
11	Detection of the <i>M1841</i> mutation in HIV DNA is suggestive of the presence of a defective genome due to the APOBEC enzyme when it is associated with other suggestive mutations (e.g., <i>M411</i> , <i>M2301</i> on reverse transcriptase) and/or stop codons.	0% (0)	9.5% (2)	90.5% (19)	8	Strong consensus	3.1% (1)	15.6% (5)	81.3% (26)	8.5	Strong consensus	STRONG CONSENSUS
12	When resistance mutations attributable to APOBEC are detected, their significance should be interpreted with caution and based on the clinical context and treatment history.	4.8% (1)	0% (0)	95.2% (20)	9	Strong consensus	5.4% (2)	5.4% (2)	89.2% (33)	9	Strong consensus	STRONG CONSENSUS
13	The presence of resistance mutations attributable to APOBEC should be reported in the HIV DNA genotyping analysis.	9.5% (2)	4.8% (1)	85.7% (18)	8	Strong consensus	N.A.	N.A.	N.A.	N.A.	N.A.	STRONG CONSENSUS
Reporting of genotyping results												
14	The clinical interpretation of resistance mutations on HIV DNA genotyping should be discussed at multidisciplinary consultation meetings.	4.8% (1)	19% (4)	76.2% (16)	9	Strong consensus	4.8% (1)	19% (4)	76.2% (16)	9	Strong consensus	STRONG CONSENSUS
15	In a virologically suppressed patient, the detection on HIV DNA genotyping of new resistance mutations (excluding APOBEC & stop codons) previously undetected must be considered for a treatment switch decision and subsequent follow-up.	4.8% (1)	14.3% (3)	81% (17)	8	Strong consensus	4.8% (1)	14.3% (3)	81% (17)	8	Strong consensus	STRONG CONSENSUS
ARV	ARV recycling											
16	In a patient with an undetectable viral load for at least one year, in the presence of a documented M184V substitution over the past five years, the use of a 2^{nd} generation InSTI + XTC + 1 NRTI combination presents low risk of virological failure over time.	14.3% (3)	9.5% (2)	76.2% (16)	8	Strong consensus	4.3% (2)	6.4% (3)	89.4% (42)	8	Strong consensus	STRONG CONSENSUS

17	In a patient with an undetectable viral load for at least one year, in the presence of a documented $M184V$ substitution over the past five years, the use of a 2 nd generation InSTI + XTC combination might present a risk of virological failure over time.	19% (4)	4.8% (1)	76.2% (16)	8	Strong consensus	22.5% (9)	25% (10)	52.5% (21)	7	No consensus	NO CONSENSUS
18	The use / recycling of NNRTIs in the event of documented resistance to ARV agents of this class is associated with a greater risk of virological failure, independently of the duration of undetectable viral load, especially in drug-reduction strategies using ARV agents of this class.	4.8% (1)	19% (4)	76.2% (16)	8	Strong consensus	0% (0)	17.9% (7)	82.1% (32)	8	Strong consensus	STRONG CONSENSUS
Test availability and delays												
19	HIV DNA genotypic tests should be accessible in clinical practice to all clinicians managing PLWH.	4.8% (1)	9.5% (2)	85.7% (18)	9	Strong consensus	10.6% (5)	6.4% (3)	83.0% (39)	9	Strong consensus	STRONG CONSENSUS
20	Reports of genotypic HIV DNA test results should be given to clinicians within 30 days.	4.8% (1)	4.8% (1)	90.5% (19)	9	Strong consensus	10.0% (4)	7.5% (3)	82.5% (33)	9	Strong consensus	STRONG CONSENSUS

N.A.: Not available; ARV: Antiretroviral; APOBEC: Apolipoproteins B mRNA editing enzyme; InSTI: Integrase Strand Transfer Inhibitor; XTC: Lamivudine/Emtricitabine; PLWH: People living with HIV

Table S3 – APOBEC context drug resistance mutations

ARV class	Major resistance mutations								
PI	M46I, G73S, D30N								
NRTI	D67N, M184I								
NNRTI	M230I, E138K, G190E, G190S								
InSTI	G163R, G163K, G140R, D232N, E138K, G140S, G118R, R263K								

PI: Protease Inhibitor; InSTI: Integrase Strand Transfer Inhibitor