



Heterogeneous indications and the need for viability assessment: An international survey on the use of machine perfusion in liver transplantation

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Abstract

Although machine perfusion (MP) is being increasingly adopted in liver transplantation, indications, timing, and modality are debated. To investigate current indications for MP a web-based Google Forms survey was launched in January 2021 and addressed to 127 experts in the field, identified among first and corresponding Authors of MP literature in the last 10 years. The survey presented 10 real-life cases of donor–recipient matching, asking whether the liver would be accepted (Q1), whether MP would be used in that particular setting (Q2) and, if so, by which MP modality (Q3) and at what timing during preservation (Q4). Respondents could also comment on each case. The agreement was evaluated using Krippendorff's alpha coefficient. Answers from 39 (30.1%) participants disclosed significant heterogeneity in graft acceptance, MP indications, technique, and timing. Agreement between respondents was generally poor (Q1, $\alpha = 0.11$; Q2, $\alpha = 0.14$; Q3, $\alpha = 0.12$, Q4, $\alpha = 0.11$). Overall, respondents preferred hypothermic MP and an end-ischemic approach in 56.3% and 81.1% of cases, respectively. A total of 18 (46.2%) participants considered only one MP approach, whereas 17 (43.6%) and 3 (7.7%) considered using alternatively 2 or 3 different techniques. Of 38 comments, 17 (44.7%) were about the use of MP for graft viability assessment before implantation. This survey shows considerable variability in MP indications, emphasizing the need to identify scenarios of optimal utilization for each technique. Viability assessment emerges as a fundamental need of transplant professionals when considering the use of MP.

KEYWORDS

ex situ organ perfusion, machine perfusion, normothermic regional perfusion, organ preservation, viability assessment



1 | INTRODUCTION

Machine perfusion (MP) is revolutionizing the field of liver transplantation (LT).^{1,2} The deep interest in MP, a term that encompasses various approaches, is supported by the impelling necessity to increase donor pool and to improve the preservation of grafts from extended criteria donors. This necessity fostered its transition from the experimental setting to the clinical practice. A rapidly growing body of literature supports its use in different scenarios in LT.³⁻²³ Three randomized controlled trials have been published,^{9,11,19} whereas several others are currently ongoing and results are expected soon.²⁴

In keeping with the relatively recent adoption, MP indications and modality appear to be highly variable among different centers. Which grafts should be preserved using MP, in which scenarios are its cost justified, what technique is to be preferred and what is the optimal timing for MP use during preservation are all open questions. Furthermore, the choice of a particular approach may be influenced by each center experience and skills, different organ procurement organizations, and health-care policies.

Despite its clinical relevance, variability in MP indications among different centers has not been investigated so far. Thus, we sought to explore different approaches to MP in LT by analyzing how some real-life donor-recipient scenarios, gathered from the experience of the promoting center, would be managed. This study was inspired by a recent survey from the Zurich group, which evidenced considerable discrepancies in the treatment of colorectal liver metastases even among experts in the field.²⁵ The working hypothesis of this survey was that indications for MP are heterogeneous, even among transplant professionals with direct MP experience and a thorough understanding of the advantages and limitations of each technique. This survey aimed to provide an accurate and representative cross-sectional picture of how different MP modalities are perceived among various groups and regions, to identify the specific areas of interest for future consensus guidelines and highlight the issues and needs of transplant professionals toward MP technology.

2 | MATERIALS AND METHODS

A web-based Google Forms survey (<https://forms.gle/2DLK3kK1EMCWewbz6>) was launched in January 2021 and addressed by email to 127 experts in the field of liver MP, identified by a systematic PubMed search of first and corresponding authors of articles about clinical liver MP in the last 10 years. In case of nonresponse after the first contact, we sent a single reminder to target experts.

The survey link was not shared on social media platforms. Geographical location of target experts was as follows: Europe, $n = 90$ (71%); North America, $n = 24$ (19%); South America, $n = 3$ (2%); Asia, $n = 10$ (8%).

The design and ethics of this survey followed the recommendations “Best Practices for Survey Research” by the American Association for Public Opinion Research (AAPOR) (<https://www.aapor.org/>). Briefly, respondents were informed about the aim of the survey and that only pooled data would be reported. Anonymity was preserved at each stage. For stratification purposes, geographical location, volume and presence of an established MP program at respondent center was asked in preliminary questions. Noteworthy, also respondents from centers without an established MP program had direct clinical experience with MP and proven knowledge in the field of MP. To safeguard anonymity, email address could be provided on a voluntary basis at a separate link (<https://forms.gle/iENmPygSYDrtoEMLA>) where respondents could discover how survey cases had actually been managed at the promoting center. The full text of the survey is available as Supporting Information. All participants included as co-authors accepted to be included and approved the final version of the manuscript. The survey was approved by the local Institutional Review Board that ruled out the necessity for further approval by the local Ethics Committee. Study procedures complied with the 2008 version of the Declaration of Helsinki (<https://www.wma.net>).

The survey was based on 10 real-world cases managed at the promoting transplant center in the period 2019-2020 (Table 1 and Supporting Information). For each case, the most relevant information available during the allocation process was provided. Data included clinical donor data, blood test results, macroscopic evaluation by the retrieving surgeon, availability of a preretrieval biopsy and, if so, histologic features like degree of macrovesicular steatosis. A brief picture of recipient clinical data and degree of urgency was also provided, to allow respondents also answering on the ground of donor-recipient matching.

Cases were chosen to reflect the heterogeneity of scenarios that may be faced in everyday practice. Particularly, three cases (cases 1, 7, and 10) were characterized by advanced donor age (>75 years). Of these, one (case 1) was further complicated by the association of significant graft macrovesicular steatosis (20%), whereas another one was characterized by graft allocation to an unstable recipient (case 7). In two cases (case 2 and 6), graft appeared steatotic, with a liver biopsy showing 40% macrovesicular steatosis (case 2). There were three cases of donation after circulatory death (DCD), one controlled (Maastricht 3, case 4) and two uncontrolled (Maastricht 2, case 3 and 9), all had prolonged warm ischemia time

**TABLE 1** Brief description of survey cases (for a full description see Supporting Information)

	Donor issues	Recipient issues	MP ¹	Transplanted	Outcome
Case 1	78-yo DBD with 20% macrosteatosis	None	NMP	Yes	Dead at 6 months due to HCC recurrence; no ITBL
Case 2	63-yo DBD with 40% macrosteatosis	MELD = 20; BMI = 33	D-HOPE	Yes	Retransplanted on POD 31st; dead at 3 months due to HHV8 infection
Case 3	49-yo type II DCD with 35-minutes asystolic WIT	MELD = 21	NRP + D-HOPE	Yes	Alive at 12 months; no ITBL
Case 4	56-yo type III DCD with 49-minutes fWIT	None	NRP + D-HOPE	Yes	Alive at 10 months; no ITBL
Case 5	52-yo DBD with elevated liver enzymes	None	D-HOPE	Yes	Alive at 14 months; no ITBL
Case 6	42-yo DBD with steatotic appearance	None	D-HOPE	Yes	Alive at 14 months; no ITBL
Case 7	76-yo DBD with elevated liver enzymes	HBV-related ALF; MELD = 41	NMP	Yes	Alive at 20 months; no ITBL
Case 8	12-yo DBD with elevated liver enzymes	None	D-HOPE	Yes	Alive at 21 months; isolated S6 duct ITBL + anastomotic stricture
Case 9	21-yo type II DCD with 31-minutes asystolic WIT	None	NRP + NMP	No	Na
Case 10	96-yo DBD	None	D-HOPE	Yes	Alive at 2.5 years; no ITBL

Abbreviations: ALF, acute liver failure; BMI, body mass index; DBD, donation after brain death; DCD, donation after circulatory death; D-HOPE, dual hypothermic oxygenated machine perfusion; fWIT, functional warm ischemia time; HBV, hepatitis B virus; ITBL, ischemic-type biliary lesion; MELD, model for end-stage liver disease; MP, machine perfusion; Na, not applicable; NMP, normothermic machine perfusion; NRP, normothermic regional perfusion; WIT, warm ischemia time.

¹Choice of machine perfusion technique at the promoting center.

and underwent normothermic regional perfusion prior to retrieval. A pediatric donor with liver enzymes suggesting acute liver injury was also included (case 8). Two cases were classified as low complexity cases (case 5 and 6), whereas all others were classified as high-complexity cases.

For each case, respondents were asked whether they would accept the offer (Q1), whether they would use MP (Q2) and, if so, which modality they would choose (Q3) and at what timing during preservation (Q4). A free comment could be added at the end of each case (Q5).

MP techniques were defined according to standard definitions.²⁶ A sequential approach was defined as hypothermic MP followed by normothermic MP, with or without interposing a phase of controlled oxygenated rewarming.¹⁷

To investigate the perception of advantages and limitations of each different technique, contributors were specified to answer based not strictly on their center practice but on what they would have done in an ideal setting in which all techniques were available, free from any funding/logistic/resource restriction.

Data are expressed as number (%) and median (interquartile range) and compared using Mann-Whitney, Chi-square and Fisher tests, as appropriate. Interrater reliability was expressed using percentage of agreement and Krippendorff's alpha coefficient, which takes into account casual agreement by chance. Values vary from 1 (perfect agreement) to -1 (perfect disagreement, exceeding what can be expected by chance). A value of 0 reflects no interrater agreement beyond casual agreement by chance. Alpha > 0.8 is generally accepted as a measure of good interrater agreement. Statistical analyses and data visualization were performed using R: A language and environment for statistical computing (R Foundation for Statistical Computing, Vienna, Austria).

3 | RESULTS

Response rate was 30.1% (39/127). Of respondents, 10 were senior figures or program directors at their Institution. Most respondents were from Europe (n = 35, 89.7%), whereas 3 (7.7%) and 1 (2.6%) were from



FIGURE 1 Survey answers [Color figure can be viewed at wileyonlinelibrary.com]

North America and South America, respectively. Centre volume was ≥ 100 , 50-100, and < 50 for 10 (25.6%), 19 (48.7%), and 10 (25.6%) respondents, respectively. Five respondents were from centers without an established MP program.

Figures 1 and 2 summarize participants' choices in each case, highlighting significant heterogeneity toward MP indication, modality, and graft acceptance. Lack of agreement between participants was confirmed by inter-rater reliability analysis, which showed Krippendorff's alpha coefficient never exceeding 0.15 for any of the survey questions. Lack of agreement persisted even when

specific subsets (elderly donors, steatotic grafts, DCD donors, high-complexity and low-complexity cases) were analyzed separately (Table 2).

Of 390 theoretical maximum times that MP could have been deemed indicated (ie, in case all respondents would have used MP in all survey cases), MP was deemed indicated 174 times (44.6%). Hypothermic MP was the most preferred approach in 98 (56.3%) cases, followed by normothermic ($n = 63$, 36.2%), sequential ($n = 10$, 5.7%), and subnormothermic MP ($n = 3$, 1.7%). One participant (2.6%) did not consider MP useful in any case, whereas 18 (46.2%), 17 (43.6%), and 3 (7.7%) considered one, two,

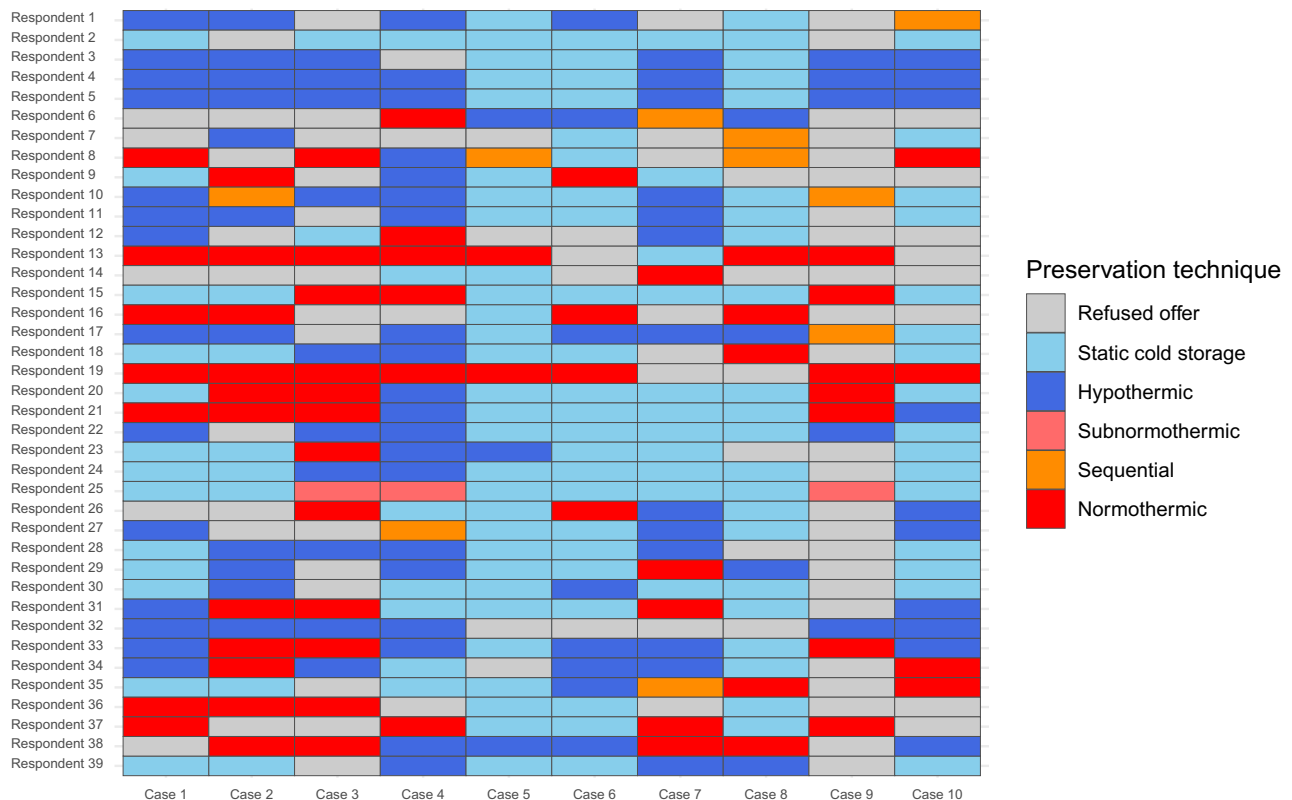


FIGURE 2 Heatmap depicting respondents' attitude with regard to graft acceptance and choice of preservation method. Sequential approach includes cases in which hypothermic machine perfusion (MP) was followed by normothermic MP, with or without interposing a phase of controlled oxygenated rewarming. Resp, respondent (followed by sequential number) [Color figure can be viewed at wileyonlinelibrary.com]

or three different MP approaches, respectively. Preference for an MP technique varied significantly according to the presence of an established MP program, with participants from centers without an MP program favoring more frequently hypothermic MP (Table 3).

When MP was indicated, an end-ischemic approach was preferred in 137 (81.1%) cases, whereas continuous perfusion throughout preservation was chosen in 32 (18.9%). In five cases, preferred timing was not indicated. The choice of the technique influenced timing, with hypothermic perfusion being associated more frequently with an end-ischemic approach and normothermic perfusion with continuous use throughout preservation (Table 4).

Finally, of 38 free comments to survey cases, 17 (44.7%) stressed the need for viability testing before transplant in cases 1, 2, 3, 4, 6, 9, and 10. Viability testing invariably involved the use of normothermic MP, either alone or in the setting of a sequential approach. Other comments were about logistic/organizational aspects ($n = 10$), the need of ruling out significant atherosclerosis of the hepatic artery ($n = 3$), the utility of obtaining a liver biopsy ($n = 2$), or other issues ($n = 6$). A full list of participants comments is available in Table S1.

4 | DISCUSSION

The results of this survey could be summarized in three main points. First, there were significant discrepancies in MP indications and technique among participants and, more surprisingly, the same heterogeneity was observed with regard to graft acceptance, which was equally highly variable. Second, while approximately half of the participants stuck to one favorite technique regardless of the peculiarities of each case, the other half proposed two or more different approaches to be adapted to different scenarios. Third, the majority of comments stressed viability assessment as a fundamental aspect of MP, which was invariably linked to the use of normothermic MP.

MP indications and techniques are strongly varying, which can be attributed to the very recent clinical reintroduction of this technology after the early experiences in the pioneering era of solid organ transplantation.²⁷ As of today, three randomized controlled trials have been published,^{9,11,19} one of which after the completion of this survey.¹⁹ Along with other retrospective studies, current literature suggests that MP conveys a significant advantage over static cold storage in several settings, including DCD donors,^{7,15,18,19} extended-criteria DBD donors,^{12,14} elderly

**TABLE 2** Agreement between respondents according to setting and complexity of cases

	All cases	Age > 75	Steatotic	DCD	Low complexity	High complexity
Q1: graft acceptance						
PA	69%	70%	73%	62%	77%	68%
α	0.11	-0.01	0.02	0.17	-0.01	0.12
CI	(-0.05, 0.27)	(-0.03, 0.02)	(-0.10, 0.14)	(-0.27, 0.62)	(-0.02, 0.01)	(-0.05, 0.29)
Q2: use of MP (yes/no)						
PA	45%	38%	43%	51%	50%	44%
α	0.14	-0.01	0.09	0.11	0.01	0.09
CI	(0.04, 0.25)	(-0.02, 0.01)	(0.07, 0.12)	(-0.15, 0.37)	(-0.04, 0.07)	(-0.01, 0.20)
Q3: MP preferred approach						
PA	33%	26%	32%	34%	46%	30%
α	0.11	-0.01	0.06	0.09	0.01	0.08
CI	(0.02, 0.20)	(-0.02, 0.01)	(-0.05, 0.18)	(-0.11, 0.30)	(-0.04, 0.05)	(-0.01, 0.16)
Q4: Timing of MP						
PA	38%	30%	37%	41%	50%	35%
α	0.12	-0.01	0.08	0.09	0.01	0.08
CI	(0.02, 0.21)	(-0.02, 0.01)	(-0.05, 0.21)	(-0.12, 0.30)	(-0.03, 0.042)	(-0.01, 0.17)

Notes: Subgroups: Age > 75, case 1, 7 and 10; Steatotic, case 2 and 6; DCD, case 3, 4 and 9; Low complexity, case 5 and 6; High complexity, case 1, 2, 3, 4, 7, 8, 9, 10.

Abbreviations: CI, confidence interval for Krippendorff's alpha coefficient; DCD, donation after circulatory death; MP, machine perfusion; PA, percentage of agreement; Q, question; α , Krippendorff's alpha coefficient.

TABLE 3 Preferred machine perfusion approach

	Overall	MP program		P^*
		No	Yes	
n	174	19	155	
MP technique				<.01
Hypothermic	98 (56.3)	12 (63.2)	86 (55.5)	
Normothermic	63 (36.2)	3 (15.8)	60 (38.7)	
Sequential approach	10 (5.7)	1 (5.3)	9 (5.8)	
Subnormothermic	3 (1.7)	3 (15.8)	0 (0.0)	

Note: Data are presented as number (%).

*Chi-square test comparing group with and without an established MP program.

TABLE 4 Timing of machine perfusion according to technique

	End-ischemic (back-to-base)	Throughout perfusion	P^{**}
Hypothermic	84 (61.3)	11 (34.4)	<.01
Normothermic	44 (32.1)	17 (53.1)	
Sequential*	9 (6.6)	1 (3.1)	
Subnormothermic	0 (0.0)	3 (9.4)	

Note: Data are expressed as number (%) and represent the aggregate number of times each timing (end-ischemic versus throughout perfusion) was chosen according to MP technique.

*Sequential approach includes cases in which hypothermic MP was followed by normothermic MP, with or without interposing a phase of controlled oxygenated rewarming; **Chi-square test.



donors⁹ and steatotic grafts.¹⁰ However, the thresholds and clinical criteria for its application are difficult to determine in the real-life practice, all the more that use of MP, at least initially, somewhat alters standard retrieval-transplant routine and increases the costs of allograft preservation.²⁸ In this context, recent guidelines proposed by the Italian Society for Organ Transplantation (SITO)²⁹ and the International Liver Transplantation Society (ILTS)^{30,31} represent a valuable effort to help clinicians evaluating MP indication in different settings and designing future trials on liver MP.

As expected, the choice of MP technique was arbitrary, subjective, or center-related. Indeed, early studies have assessed feasibility of MP in a clinical setting or compared MP with SCS. No study comparing different MP techniques head-to-head has been published so far.²⁸ However, results of a large, multicenter study comparing end-ischemic normothermic versus end-ischemic hypothermic oxygenated MP from Germany are expected and will hopefully shed some light on the respective benefits and shortcomings of each technique.

In this survey, hypothermic MP was chosen in 56.3% of cases, and an end-ischemic approach was by far the most preferred timing (81.1% of cases). The reason why most respondents tended to prefer hypothermic MP might be at least partially related to the fact that hypothermic MP is frequently perceived as technically easier to perform and associated with inferior technology-related risks. Noteworthy, results of the DHOPE-DCD trial,¹⁹ favoring the use of hypothermic MP over static cold storage, were published after the completion of this survey.

Concerning MP timing, the vast preference toward an end-ischemic approach likely reflects the perceived logistical challenges linked to backtable preparation and MP setup at the retrieval hospital and the risks associated with potential device malfunction during organ transport. However, it is worth noticing that current evidence^{11,32} does not support the concept that preservation by upfront MP is associated with increased risk.

An interesting and rather unexpected finding was that half of the respondents proposed more than one technique to be applied in different scenarios. As any MP technique requires a certain degree of expertise, we would have expected that most respondents would feel more comfortable with one particular technique. In contrast, half of the respondents chose at least two different techniques for the proposed scenarios, suggesting that different techniques were perceived as associated with distinct characteristics and goals. Some respondents considered some techniques not as competitive, but rather as complementary, such as end-ischemic hypothermic MP for initial resuscitation of the graft, followed by controlled oxygenated rewarming and subsequent normothermic MP for viability assessment.¹⁷

Finally, viability assessment was frequently indicated as a unique possibility offered by MP. In this regard, MP has radically changed the concept of organ preservation, creating a time window during which a functioning organ can be inspected and evaluated *ex situ*. This concept has already allowed successfully recovering and transplanting organs previously deemed unsuitable for LT.^{17,20,33} In this survey, viability assessment were invariably associated with the use of normothermic MP. However, as brilliantly summarized by Brüggewirth et al,^{34,35} viability assessment is still an imperfect science, as many criteria have been proposed but none validated. Many criteria focus on hepatocyte function, whereas cholangiocyte function and injury have been rather neglected.^{2,36} Furthermore, some promising findings from the Zurich group suggest that viability assessment is possible also during hypothermic MP, although validation of this claim is still lacking.³⁷ Ultimately, viability assessment appears to be a fundamental need of the clinician toward both hypothermic and normothermic MP technologies, especially in cases characterized by significant uncertainty regarding the suitability and function of the allograft.

Taken together, these findings depict a picture of non-uniform MP practice among experts in the field and highlight the urgent need for more clinical data and studies to make an evidence-based approach to MP utilization possible. The recently released ILTS consensus guidelines on *ex situ* liver MP have addressed relevant aspects and highlighted potential pitfalls of future MP trials, tracing the path toward a more solid evidence in the field.^{30,31} As proposed, a necessary step in this direction would be wide and transparent data sharing by creation of registries detailing indications and outcome of MP cases. Future studies should focus on clinically relevant endpoints, whereas recently introduced scores³⁸⁻⁴⁰ could be used as measures of post-LT graft function.

This study has limitations, including limited sample size ($n = 39$) and response rate (30.1%), although this last was in line with that of other web-based medical surveys.⁴¹ As the majority of participants were from Europe, results do not accurately reflect practices elsewhere. Significant differences in donation rate, waiting list time, surgeons personal experience and country-specific protocols, especially with regards to DCD donors (systematic use of end-ischemic hypothermic MP vs. normothermic regional perfusion), could have influenced variance in answers. As an example, use of normothermic regional perfusion in the proposed DCD cases was obligatory at the promoting center, and this is why this was not considered as a further preservation option. However, normothermic regional perfusion undoubtedly represents another variable in the equation and its use, alone or in association with MP, certainly deserves evaluation.^{8,42}



One strength of this survey was its being targeted to people with direct hands-on experience with MP and with a solid background about the advantages and limitations of each technique.

In conclusion, just like what happens when evaluating an organ offer, it appears that the choice of applying MP is based on an important element of subjectivity and regional differences.⁴³ Every clinician likely takes into account several donor, recipient and logistic factors, which are weighed based on her/his experience, retrieval organization and waitlist pressure. This highlights the need for high-quality studies, focusing on clearly defined settings and having strong clinical endpoints, to drive clinical MP use in everyday practice.

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CONFLICT OF INTEREST

The authors declare that they have no conflicts of interest with the contents of this article.

AUTHORS' CONTRIBUTIONS

DP: concept and design, data collection, analysis and interpretation, and drafting the article; DC, FR: concept and design, and data collection; RA, MIB, EB-R, IMAB, ZC, RDC, VEDM, DD, DE, DG, AJH, DK, QL, GL, TMM, AM, FM, DN, AN, DP, MR, MCS, AW, PDS, CF, WJ, MM, RJP, MR, MS, MS: data interpretation, critical revision of article, and approval of the article; AWA: concept and design, data interpretation, critical revision of article, and approval of article; RR: data interpretation, critical revision of article, and approval of the article.

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SUPPORTING INFORMATION

Additional Supporting Information may be found online in the Supporting Information section.

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**APPENDIX A**

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