Survey of transfusion practices in preterm infants in Europe

Alexandra Scrivens 1, Nora Johanna Reibel 2, Lisanne Heeger,3,4 Simon Stanworth,5 Enrico Lopriore 6, Helen V New,6 Christof Dame,7 Karin Fijnvandraat,8 Emöke Deschmann,9 Marta Aguas,10 Kristin Bække,11 Francesco Stefano Cardona 12, Filip Cools,13 Ryan Farrugia,14 Stefano Ghirardello,15 Jana Lozar,16 Katarina Matasova,17 Tobias Muehlbacher,18 Ulla Sanklamp,19 Henrique Soares,20 Miklos Szabo,21 Tomasz Szczapa,22 Gabriela Zaharie,23 Charles Christoph Roehr 24,25 Suzanne Fustolo-Gunnink,24,26,27 On behalf of the Neonatal Transfusion Network

ABSTRACT

Background Preterm infants commonly receive red blood cell (RBC), platelet and fresh frozen plasma (FFP) transfusions. The aim of this Neonatal Transfusion Network survey was to describe current transfusion practices in Europe and to compare our findings to three recent randomised controlled trials to understand how clinical practice relates to the trial data.

Methods From October to December 2020, we performed an online survey among 597 neonatal intensive care units (NICUs) caring for infants with a gestational age (GA) of <32 weeks in 18 European countries.

Results Responses from 343 NICUs (response rate: 57%) are presented and showed substantial variation in clinical practice. For RBC transfusions, 70% of NICUs transfused at thresholds above the restrictive thresholds tested in the recent trials and 22% below the restrictive thresholds. For platelet transfusions, 57% of NICUs transfused at platelet count thresholds above 25×109/L in non-bleeding infants of GA of <28 weeks, while the 25×109/L threshold was associated with a lower risk of harm in a recent trial. FFP transfusions were administered for coagulopathy without active bleeding in 39% and for hypotension in 25% of NICUs. Transfusion volume, duration and rate varied by factors up to several folds between NICUs.

Conclusions Transfusion thresholds and aspects of administration vary widely across European NICUs. In general, transfusion thresholds used tend to be more liberal compared with data from recent trials supporting the use of more restrictive thresholds. Further research is needed to identify the barriers and enablers to incorporation of recent trial findings into neonatal transfusion practice.

INTRODUCTION

Blood component transfusions of red blood cells (RBCs), platelets and fresh frozen plasma (FFP) are commonly administered to preterm infants, but the evidence base for these transfusions, particularly for platelets, has thus far been limited.1 Since 2019, three large randomised controlled trials (RCTs) were published. The Effects of Transfusion Thresholds on Neurocognitive Outcomes of Extremely Low-Birth-Weight Infants (ETTNO) and...
Transfusion of Prematures (TOP) trials reported no difference between the effects of liberal versus restrictive RBC thresholds on death or neurocognitive deficit at 2 years' corrected age. 2-4 The Platelets for Neonatal Transfusion 2/Management of Thrombocytopenia in Special Subgroup: Neonates (PlaNeT-2/MATISSE) trial compared liberal (50×10^9/L) versus restrictive (25×10^9/L) platelet count thresholds, reporting a higher rate of death and major bleeding in the liberal platelet transfusion threshold group (26% vs 19%). 5 This effect was shown to be present irrespective of varying baseline risk of outcome. 6 The extent to which findings of the aforementioned trials correspond with clinical practice in Europe is unknown. The aims of the study were to describe current transfusion practices and to compare these to the recently generated evidence from clinical trials.

METHODS
This survey was performed by the Neonatal Transfusion Network (NTN) (www.neonataltransfusionnetwork.com), an international research group which aims to generate evidence to improve clinical practice in neonatal transfusion medicine. An NTN panel of four neonatologists (EL, ED, CD and CCR), one trainee neonatologist (AS), three haematologists (SJS, HN and KF) with paediatric transfusion expertise, and one clinical epidemiologist (SFFG) developed a preliminary list of questions. Topics included RBC, platelet and FFP transfusion practices in premature neonates of less than 32 weeks' gestational age (GA) at birth, addressing transfusion thresholds or indications, durations and volumes, concomitant use of diuretics, withholding enteral feeding and parental consent. We used a ranking procedure to obtain a final set of 31 questions, which we entered into LimeSurvey (LimeSurvey GmbH, Hamburg, Germany) (online supplemental materials). Neonatologists from 18 European countries volunteered to disseminate the survey. These national coordinators received a password protected link to the questionnaire, which they disseminated between October and December 2020 to neonatal care intens unit (NICUs) providing care for infants born at <32 weeks of GA. In the United Kingdom only larger regional NICUs were approached for participation, as these are known to dictate local transfusion practices. We limited responses to one per NICU. National coordinators were free to use their own contacts or use an existing neonatal network (online supplemental materials).

We extracted the LimeSurvey data to SPSS V.27, for data cleaning and analysis, by two authors working independently. We used GraphPad Prism V9.0.1 for Windows (GraphPad Software, San Diego, California USA) for graphs. We excluded confirmed double entries, ineligible responses and responses that were >75% blank, and converted haematocrit to haemoglobin using this formula: haemoglobin (g/L) = haematocrit(%) × 300.

The TOP and ETTN0 trial had only recently been published at the moment of survey dissemination. 2,3 Therefore, we did not aim to assess implementation of their results but instead to assess how current clinical practice compared with the thresholds tested in these trials. To make the comparison, we combined the two trials to select one liberal and one restrictive 'ETTN0/TOP threshold' for each of our 15 survey clinical scenarios: ‘air’, ‘low flow’, ‘high flow’ (</>30% FiO2) and ‘intubated’ for three postnatal age groups. Where ETTN0 and TOP thresholds differed, we selected the higher liberal value and the lower restrictive. The low flow and high flow’ clinical scenarios could not be assigned to either the ‘critical’ or the ‘non-critical’ strategies in the ETTN0 and TOP trials because of overlapping definitions. Therefore, some clinical scenarios were classified as both critical and non-critical in one or both trials, leading to relatively wide ETTN0/TOP threshold ranges (online supplemental tables S1 and S2). We calculated the proportion of reported thresholds in our survey that were at or above the liberal ETTN0/TOP threshold, between the liberal and restrictive ETTN0/TOP threshold or at or below the restrictive ETTN0/TOP threshold, for all clinical scenarios for both RBC and platelet transfusion thresholds.

We performed two sensitivity analyses. To estimate the effect of non-responder bias, we compared early versus late responders. This is an established method to estimate if responders answered survey questions differently from non-responders, where late responders are considered a proxy for non-responders. We combined the first and last 20% of entries in each country to form the early-responder and late-responder groups. We also assessed the effect of varying response rates between countries in a weighted sensitivity analysis, where entries received a weight equal to one divided by the response rate in their respective country.

RESULTS
Response rate
After removal of seven duplicate responses, 10 ineligible responses and 168 responses which were >75% blank, responses from 343 NICUs were included, yielding an overall response rate of 57% (343/597). The response rate per country varied between 21% and 100% (median 81%). We included NICUs in Austria (7 of a total of 7 units), Belgium (16/19), Finland (5/5), Germany (112/160), Hungary (21/21), Italy (49/105), Malta (1/1), the Netherlands (9/9), Norway (6/6), Poland (18/36), Portugal (10/11), Romania (4/19), Slovakia (10/13), Slovenia (3/4), Spain (41/111), Sweden (8/8), Switzerland (6/8) and the United Kingdom (17/55). UK

RBC transfusion thresholds
A violin plot of haemoglobin thresholds for RBC transfusions is shown in figure 1A. A total of 104 NICUs used haematocrit thresholds, which were converted into haemoglobin. Higher haemoglobin thresholds were adopted as the degree of respiratory support intensified. On average, over all 15 clinical scenarios, 22% of reported thresholds were below, and 8% at the restrictive ETTN0/TOP threshold, 68% were in between the restrictive and liberal ETTN0/TOP threshold, <1% at the liberal threshold and 2% above the liberal ETTN0/TOP threshold (figure 1B).

Platelet transfusion thresholds
A violin plot of platelet count thresholds for platelet transfusions is shown in figure 2. Platelet transfusion thresholds above 25×10^9/L were used in 57% (188/332) and 47% (158/333) of NICUs for a non-bleeding infant of <28 weeks' GA or 28–32 weeks' GA, respectively. For infants treated with ibuprofen, platelet transfusion thresholds above 25×10^9/L were used in 84% (249/297 and 248/297) of NICUs for infants of <28 weeks' GA and infants of 28–32 weeks’ GA, respectively. For infants treated with ibuprofen, platelet transfusion thresholds above 25×10^9/L were used in 84% (249/297 and 248/297) of NICUs for infants of <28 weeks' GA and infants of 28–32 weeks’ GA. Thresholds of 20×10^9/L or less were used in 27% (91/332) for infants with GA of <28 weeks without bleeding, 34% (114/333) for infants with GA of 28–32 weeks without bleeding, 8% (25/297) for infants with GA of <28 weeks and ibuprofen and 9% (26/297) of infants with GA of 28–32 weeks with ibuprofen. Infants with lumbar puncture, surgery or active bleeding could not be compared with the trial results as they were allowed additional transfusions at the
Figure 1  (A) Violin plots of haemoglobin-based RBC transfusion thresholds for different postnatal age and respiratory support levels in infants born at 30%=infants on 30% oxygen by non-invasive respiratory support (including continuous positive airway pressure, biphasic intermittent positive airway pressure (synchronised or unsynchronised) and nasal high flow). Ventilated=infants who are intubated and ventilated. The grey boxes highlight the values between the restrictive and liberal ETTNO/TOP thresholds (online supplemental table S1). Cochrane SR refers to the restrictive thresholds of previous trials, summarised in a Cochrane systematic review by Whyte et al. Number of datapoints per violin plot: 325, 323, 324, 326, 327, 327, 326, 325, 326, 327, 327, 327, 326, 325 and 326 (from left to right). Violin plots are a combination of a boxplot (showing median and IQRs) with a kernel density plot (showing the distribution of the data). The wider the plot, the more NICUs selected this threshold. (B) 100% stacked bar chart showing for each clinical scenario the proportion of the reported transfusion thresholds (from top to bottom) that were above the liberal ETTNO/TOP threshold, at the liberal ETTNO/TOP threshold, between the liberal and restrictive ETTNO/TOP threshold, at the restrictive ETTNO/TOP threshold or below the restrictive ETTNO/TOP threshold. ETTNO, Effects of Transfusion Thresholds on Neurocognitive Outcomes of Extremely Low-Birth-Weight Infants; NICU, neonatal intensive care unit; RBC, red blood cell; TOP, Transfusion of Prematures.
discretion of the treating neonatologist according to the RCT protocol in the PlaNeT-2/MATISSE trial.

Fresh frozen plasma
Eleven percent of NICUs (38/332) performed routine coagulation tests on infants born at <32 weeks’ GA. FFP was used for the following indications: coagulopathy with active bleeding, 93% (320/343); active bleeding without coagulopathy, 46% (158/343); coagulopathy without active bleeding, 39% (133/343); sepsis, 26.5% (91/343); and hypotension, 25% (85/343).

Duration, volume and rates of transfusion
Transfusion volume and duration are depicted in figure 3. We calculated transfusion rates in millilitre per kilogram per hour by dividing volume (mL/kg) by duration (hours). Volumes ranged between 10 mL/kg and 20 mL/kg in 99% of transfusions. The median volume was 15 mL/kg for RBC (IQR 15–20), platelet transfusions (IQR 15–15) and FFP (IQR 15–20). Duration ranged between 1 hour and 7 hours for RBC transfusions (median 4, IQR 3–4), 15 min and 4 hours for platelets (median 1, IQR 0.5–2.0), and 30 min and 4 hours for FFP (median 2, IQR 1–3). Transfusion rates varied between 3.3 mL/kg/hour and 15.0 mL/kg/hour for RBCs (median 4.0, IQR 3.8–5.0), 3.3 mL/kg/hour and 60.0 mL/kg/hour for platelets (excluding two outliers at 120 and 80 mL/kg/hour) (median 15.0, IQR 7.5–20.0) and 2.5 mL/kg/hour and 50.0 mL/kg/hour for FFP (median 10.0, IQR 5–15).

Diuretics, enteral feeding and parental consent
Diuretics were ‘always’ or ‘sometimes’ prescribed in conjunction with RBC, platelet and FFP transfusions in 47% (154/329), 18% (57/322) and 28% (92/323) of NICUs, respectively. Enteral feeding was always or sometimes withheld during RBC transfusion in 28% (94/337) and 9% (31/337) of NICUs, respectively. Parental consenting for non-emergency transfusion practices varied, with 8% of NICUs (28/335) requiring no consent, 9% (31/335) requiring verbal consent only, 6% (20/335) requiring verbal consent documented by a clinician, 70% (241/335) requiring verbal and written consent and 4% (15/335) used other forms of consent, not otherwise specified.

Sensitivity analyses
Our weighted analysis showed no substantial changes compared with our primary analysis, suggesting that bias because of varying response rates between countries was likely limited (online supplemental tables S3–S6). The results of our non-responder analysis indicated that non-response bias may have resulted in an underestimation of platelet transfusion thresholds and the proportion of NICUs giving FFP for indications other than active bleeding, as these were higher in the late responders (online supplemental table S7).

DISCUSSION
Despite recent evidence from clinical trials, we demonstrated substantial variation in transfusion practices for RBC, platelets and FFP, regarding thresholds, volume and rate of transfusion across European NICUs. To our knowledge, this is the first survey to assess neonatal transfusion practices across Europe. Other surveys included only selected countries or included mostly non-European NICUs. A retrospective cohort study of North American NICU blood component transfusion thresholds for all infants between 2013 and 2016 by Patel et al also found wide variation in practice with regard to transfusion thresholds.

Comparison with RBC trials
The large variation in RBC transfusion thresholds probably reflects the lack of international and European consensus in transfusion criteria, and the lack of evidence until recently. Prior to 2020, clinical practice was partly based on two RCTs (the Premature Infants in Need of Transfusion and Iowa trials), which have been summarised in systematic reviews and national guidelines supporting the use of restrictive thresholds. These thresholds were roughly similar to those tested in the TOP and ETNO trials (online supplemental table S2). In our survey, reported use of thresholds similar to or above the previously tested liberal thresholds was rather rare. However the majority

Figure 2 Violin plots of platelet count transfusion thresholds for different clinical scenarios. Datapoints per violin plot: 332, 297, 316, 311, 329, 333, 297, 317, 311 and 331, from left to right. Violin plots are a combination of a boxplot (showing median and IQRs) with a kernel density plot (showing the distribution of the data). The wider the plot, the more NICUs selected this threshold. MATISSE, Management of Thrombocytopenia in Special Subgroup.
(70%) of thresholds reported were above the TOP/ETTNO restrictive thresholds, while surprisingly, 22% reported thresholds below the restrictive thresholds. These results could be due to implementation of evidence from earlier trials but could also indicate that neonatologists are uncertain regarding restrictive thresholds. Arguments against restrictive thresholds include the possibility of minor neurodevelopmental abnormalities or impairments that become apparent at a later age, which have not been assessed in existing trials. Arguments against a liberal threshold include the lack of clinical benefit of liberal thresholds in clinical trials, and observational studies suggesting potential transfusion-related harm. The reasons for the use of such low haemoglobin thresholds in 22% of units are unclear but may represent an underestimation of transfusion threshold, as we had to use wide ranges for several clinical scenarios as described previously (online supplemental table S1). Without further clinical trials, it is not known whether these low haemoglobin thresholds are safe for neonates, as various preclinical studies have suggested adverse outcomes following prolonged severe anaemia.31–33

Comparison with platelet transfusion trials
In our survey, 47%–57% of NICUs indicated using platelet count thresholds above $25 \times 10^9$/L for stable non-bleeding infants. Given the strong and concerning evidence for platelet transfusion-mediated harm, the use of platelet transfusion thresholds above $25 \times 10^9$/L for non-bleeding neonates should be discouraged while we await results from long-term neurodevelopmental analyses. Our study also showed that 27%–34% of NICUs use thresholds equal to or lower than $20 \times 10^9$/L for non-bleeding infants. The use of thresholds lower than those tested in trials may be an acceptable practice for platelet transfusions, given the evidence in favour of the restrictive versus liberal threshold, but again this needs to be tested in a clinical trial. Possible explanations for platelet transfusion-mediated adverse effects include the role of platelets in inflammatory and immunoregulatory responses, a developmental mismatch between adult donor platelets and neonatal platelets and volume-mediated effects, since platelet transfusions are given at relatively high volumes and rates compared with adult transfusions.

Fresh frozen plasma
There are no recent trials investigating FFP transfusion indications, but based on observational and adult data, most guidelines recommend that FFP should not be administered to non-bleeding infants to correct abnormalities of the coagulation screen alone.13–30 Minor coagulation abnormalities are poor predictors of bleeding risk and FFP administration often will not correct these abnormalities.30 In addition, they can be difficult to interpret, particularly for very preterm babies. We found that 39% of NICUs in our survey administered FFP in case of coagulopathy without bleeding. Furthermore, 25% of NICUs transfused FFP to treat hypotension, for which there is no robust evidence, yet there are potential risks such as transfusion-related acute long injury and severe allergic reactions.32 These findings indicate an urgent need for clinical trials in this area.

Volume, duration and rate of transfusion
Transfusion duration and rate were marked in their variation. The cause for this variation is unclear, but evidence to guide practice is lacking. On average, volumes were comparable to those used in the recent trials (15 mL/kg for PlaNeT-1/2/MATISSE and TOP, 20 mL/kg for ETTNO). Only a few small studies have assessed neonatal transfusion volume and suggested that these volumes are tolerated, though there is also evidence for harm.14 15 18–20 Duration or rates of transfusions were not reported in the ETTNO and TOP trials. High-volume or high-rate transfusions may increase the risk of transfusion associated circulatory overload (TACO), a leading cause of transfusion-associated morbidity and mortality in adults.33 Neonatal TACO is poorly defined and the true incidence of this and other transfusion-related lung injuries in neonates is not known; observational studies have shown variable outcomes.37–40 42 However, it should be noted that weight-related volumes transfused to non-bleeding neonates (usually 10–20 mL/kg) are commonly higher than those for adults (typically, 350 mL for packed red cells and 200–300 mL for platelets,
which equates to <5 mL/kg for an 80 kg adult). Moreover, the vulnerability of the cerebral vasculature and increased risk of intraventricular haemorrhage in preterm infants should also be considered. Further high-quality research is urgently needed to optimise transfusion rates and volumes.

Limitations and strengths
There are some limitations to our survey. First, we requested one response per neonatal unit, which may mask intranunit variation. Second, we did not ask for detailed transfusion product information because we anticipated that these data were not always available to neonatologists. Third, our eligibility criteria were broad; we only defined that NICUs should care for neonates born at <32 weeks’ GA. We chose this pragmatic approach to be able to paint a picture of current transfusion practices across Europe and because NICU level definitions differ per country. Lastly, combining the ETNTO and TOP trials was not always possible due to varying clinical definitions, highlighting the need for uniform transfusion indications in future trials. Strengths of our survey include the relatively high response rate despite the ongoing pandemic, use of sensitivity analyses and the wide range of countries included. This is, to our knowledge, the first study to map European neonatal transfusion practices, and given the recently published trials, our data are timely and provide a valuable starting point for further studies and highlight the need for guideline development.

CONCLUSIONS
In Europe, transfusion practices for preterm infants vary widely. Transfusion thresholds tend to be more liberal compared with data from recent trials supporting the use of more restrictive thresholds. Transfusion indications, volume, duration, concomitant use of diuretics, withholding enteral feeds and parental consent requirements differ considerably. These areas, including factors affecting the implementation of research findings, deserve further attention and clinical research.

Author affiliations
1Newborn Care Unit, Oxford University Hospitals NHS Foundation Trust, Oxford, UK
2Neonatology, Charité Universitätsmedizin Berlin, Berlin, Germany
3Neonatology, Leiden University Medical Centre, Leiden, The Netherlands
4Sanquin Blood Supply Foundation, Amsterdam, The Netherlands
5Department of Haematology, National Health Service, Blood and Transplant, Oxford University Hospitals NHS Foundation Trust, Oxford, UK
6Paediatric Transfusion Medicine, National Health Service, Blood and Transplant, London, UK
7Neonatology, Charité – Universitätsmedizin Berlin, Berlin, Germany
8Pediatrics, Emma Children’s Hospital, Pediatric Hematology, University of Amsterdam, Amsterdam, The Netherlands
9University Hospital, Stockholm, Karolinska Institute, Stockholm, Sweden
10Servicio de Neonatología, University & Polytechnic Hospital La Fe, Valencia, Spain
11Women and Children’s division, Department of Neonatal Intensive Care, Ullevål, Oslo University Hospital, Oslo, Norway
12Department of Pediatrics and Adolescent Medicine, Division of Neonatology, Intensive Care and Pediatric Neurology, Medical University of Vienna, Wien, Austria
13Universitet Ziekenhuis Brussel, Vrije Universiteit Brussel, Brussels, Belgium
14Pediatrics, Mater Dei Hospital, Moida, Malta
15Neonatal Intensive Care and Neonatology Unit, Department of Pediatrics, Fondazione IRCCS Policlinico San Matteo, Pavia, Italy
16Neonatology, University Medical Centre Ljubljana, Ljubljana, Slovenia
17Jessenius Faculty of Medicine in Martin, University Hospital Martin, Martin, Slovakia
18Neonatology, University Hospital Zurich, Zurich, Switzerland
19Department of Pediatrics, Kuopio University Hospital and University of Eastern Finland, Kuopio, Finland
20Neonatology, Centro Hospitalar Universitário de São João, Porto, Portugal
21Division of Neonatology 1st Department of Pediatrics, Semmelweis University, Budapest, Hungary
22Department of Neonatology, Neonatal Biophysical Monitoring and Cardiopulmonary Therapies Research Unit, Poznan University of Medical Sciences, Poznan, Poland
23Neonatology, University of Medicine and Pharmacy Iuliu Hatieganu Cluj, Cluj Napoca, Romania
24National Perinatal Epidemiology Unit, Clinical Trials Unit, Oxford Population Health, Medical Sciences Division, University of Oxford, Oxford, UK
25Women and Children’s, Neonatal Intensive Care Unit, Southwest Hospital, North Bristol NHS Trust, Westonbirt on Trym, Bristol, UK
26Department of Epidemiology, Leiden University Medical Center, Leiden, The Netherlands
27Pediatric Hematology, Amsterdam University Medical Center, Amsterdam, Netherlands
28Department of Neonatology, University Hospital Zurich, Zurich, Switzerland

Twitter Alexandra Scrivens @dralexscrivens and Francesco Stefano Cardona @fracardo

Acknowledgements The authors thank Laura Moschino, Camilla Caram-Deldeler, the neonatal networks that were involved, Ann Kennedy, the National Perinatal Epidemiology Unit, University of Oxford, and all neonatologists who took the time to fill in the survey.

Collaborators The study was conducted by the executive council members and other members of the Neonatal Transfusion Network (NTN). The NTN is an interdisciplinary, international research network focused on optimising neonatal transfusion practices and research worldwide. It is governed by a steering board and includes over 130 members from over 35 countries. The network is endorsed by the European Society for Pediatric Blood and Blood Transfusion, the European Blood Alliance, the European Foundation for the Care of Newborn Infants and the International Hemovigilance Network (www.neonataltransfusionnetwork.com).

Contributors The study was planned and executed by the members of the Neonatal Transfusion Network steering committee (CCR, CD, ED, EL, HVN, JvB8, KF, SS, SFG, AS, HS, LH, NH and NfR) with significant input from MAC, KB, FSC, FC, RF, SGh, JL, KM, TM, US, HS, MS, TS and GZ, who also facilitated region-specific data collection. S-FG is guarantor.

Funding The study was in part funded by the generous support and grant (PPCC21-08/2588 and RES/02624) from Sanquin Blood Supply Foundation, Amsterdam, the Netherlands, and a postdoctoral research grant (RGP2020-09/PDRG-02/04) from The European Society for Pediatric Transfusion, Geneva, Switzerland.

Competing interests None declared.

Patient consent for publication Not applicable.

Ethics approval Ethical approval was not required as no patient-specific data were collected.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request. Data are available upon reasonable request by email to the corresponding and the senior author.

Supplemental material This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to localised terms, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iDs
Alexandra Scrivens http://orcid.org/0000-0002-3429-8007
Nora Johanna Reibelt http://orcid.org/0000-0002-6412-4606
Enrico Lopriore http://orcid.org/0000-0002-3513-5066
Francesco Stefano Cardona http://orcid.org/0000-0002-9993-617X
Tobias Muehlbacher http://orcid.org/0000-0001-8661-9645
Charles Christoph Roehr http://orcid.org/0000-0001-7965-4637

REFERENCES


