Home therapy for inherited bleeding disorders in South Africa: Results of a modified Delphi consensus process

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Background. Optimal care of patients with inherited bleeding disorders requires that bleeding episodes are treated early, or still better prevented, through extension of patient care beyond hospital-based treatment to home-based therapy. In South Africa (SA), adoption of home therapy is variable, in part owing to lack of consensus among healthcare providers on what constitutes home therapy, which patients should be candidates for it, how it should be monitored, and what the barriers to home therapy are.

Objectives. To conduct a modified Delphi process in order to establish consensus on home therapy among haemophilia healthcare providers in SA.

Methods. Treaters experienced in haemophilia care were invited to participate in a consensus-seeking process conducted in three rounds. In round 1, provisional statements around home therapy were formulated as questions and collated in a structured list. In rounds 2 and 3, evolving versions of the questionnaire were administered to participants. Consensus was defined as ≥70% agreement among the participants.

Results. The panel composition included an equal number of physicians and non-physicians. The participation rate was 100% through all three consensus rounds. The group reached consensus for 92% of the statements. Consensus of 100% was reached on starting home therapy for patients with haemophilia who are candidates for it, whether given episodically or prophylactically, may be delayed while patients are transported to healthcare facilities. This delay has been shown to result in complications of bleeds, which become more difficult to manage, delayed in their resolution, and associated with poor outcomes. Once diagnosed, haemophilia imposes a high disease and treatment burden on patients, caregivers, healthcare providers and the healthcare facilities involved in management. The disease burden includes unpredictable spontaneous acute bleeds as well as chronic complications of bleeds into the musculoskeletal system, muscle and soft tissues, which may be life or organ threatening or disabling. The treatment burden is in part due to lifelong treatment for spontaneous and traumatic bleeds with clotting factor concentrate (CFC) replacement therapy, administered by intravenous infusion and therefore requiring specific care processes and skills. Current management of bleeds in haemophilia is by intravenous replacement of the deficient clotting factor to treat or prevent bleeding episodes. Episodic treatment, the reactive modality, is associated with progressive deterioration of musculoskeletal structures, musculoskeletal function and quality of life. The proactive alternative is prophylaxis, the standard of care in haemophilia, in which patients receive regular infusions of clotting factor to prevent bleed occurrence. Replacement therapy, whether given episodically or prophylactically, may be delayed while patients are transported to healthcare facilities. This delay has been shown to result in complications of bleeds, which become more difficult to manage, delayed in their resolution, and associated with poor outcomes.
poor musculoskeletal and quality-of-life outcomes.\textsuperscript{13-17} The delay
in intravenous therapy can be mitigated by teaching patients how to
practise intravenous infusion in their home settings. Indeed, in many
comprehensive treatment centres in SA, patients are taught self-
infusion at an early age.\textsuperscript{18} It has now been shown in many studies
that intravenous infusion skill combined with early replacement therapy
has the potential to limit bleed-induced musculoskeletal damage,
limit the extent of blood loss, and improve bleed resolution and
quality of life of haemophilia patients.\textsuperscript{19-21} Recent data also indicate
that early home-based therapy is cost-effective, and is associated with
reduced pain and disability, decreased school and work absenteeism,
and reduced hospitalisation rates.\textsuperscript{22-24} Consequently, in many parts of
the world, haemophilia patients are allowed to take CFC at home in
order to infuse themselves prophylactically or to self-administer early
during an acute bleeding episode.

In a survey performed in 10 haemophilia treatment centres in
SA, the practice of home therapy varied from 0% to 70%. Part of
the explanation for this wide variability in practice is that there
was no consensus among the healthcare providers on a number of
questions on home therapy-related issues for patients with inherited
bleeding disorders. The unanswered questions include: (i) who
should be allowed to take CFC at home in order to self-infuse; (ii) what
the barriers to home therapy would be; (iii) what the criteria for
identifying patients for home therapy should be; (iv) what the
characteristics of products for use in the home setting should be;
(v) what kind of support would be required for patients on home
therapy; and (vi) how the effectiveness of home therapy should be
monitored.

The Delphi approach is an established method for consensus building
by using a series of reiterated questionnaires to facilitate a panel of
selected experts to reach agreement on a particular subject.\textsuperscript{22,23} The Delphi
approach differs from standard surveys by asking the question as
what should\textit{would} be, as opposed to surveys, which ask the question as
what\textit{is}.\textsuperscript{24} It is particularly appropriate to use the Delphi method in rare
diseases such as haemophilia for which prospectively generated data
might take time or be difficult to collect, and particularly for questions
that would be impractical to answer via other forms of research.\textsuperscript{25}
The Delphi method has frequently been used in haemophilia to reach
expert consensus on a variety of issues.\textsuperscript{26-32}

\section*{Objectives}
This study used a modified Delphi method to seek consensus among
haemophilia experts on key factors required for a successful home
therapy programme for patients with inherited bleeding disorders
in SA.

\section*{Methods}
\subsection*{Study participants}
Invited participants had expertise and knowledge in the care of
patients with haemophilia and included medical practitioners, nurse
practitioners and allied healthcare workers practising in haemophilia.
They were representative of the haemophilia fraternity in SA in terms
of expertise, experience, haemophilia population they cared for, and
healthcare professions.

\subsection*{Study design and interventions}
The Delphi method was administered by an independent facilitator
in three rounds.

\subsection*{Round 1}
In a face-to-face meeting, the objective and procedures were
outlined to all participants. Participants were assigned to two groups
and tasked to identify all haemophilia patient-, home setting-,
healthcare provider-, treatment centre- and healthcare system-related
factors involved in implementation of a successful home therapy
programme. Participants considered each factor identified and the
rationale for its inclusion. They were not allowed to give individual
opinions, recommendations or consensus at this stage. All inputs
were examined by the whole group in order to clarify possible
ambiguous statements and remove duplicates.

The resultant list of issues on haemophilia home therapy was then formulated into questions, which were captured in a study
questionnaire. The questionnaire was checked by a small core group
of participants for completeness, consistency and clarity before being
finalised. Care was taken that the statements were not rated or ranked
at this stage.

\subsection*{Round 2}
The finalised questionnaire was distributed to all participants through
an online tool provided by McMaster University in Canada. Each
participant received an invitation to complete the online questionnaire
individually. Participants were not allowed to confer or discuss their
input with each other. The timeline for completion of the survey was a
maximum of 5 days after the first notification. At the end of 5 days, all
responses were centrally collated, analysed and interpreted.

\subsection*{Round 3}
Consensus statements from round 2 were used to proceed to
round 3. All participants were asked to rank the statements in order
of importance. The results were collated and checked for the level
of agreement or disagreement among the participants.

Consensus was defined as \(\geq 70\%\) of the participants agreeing
or disagreeing on a statement. Once consensus was reached, the
statements were ranked from most to least important based on the
rank received. The participant responses were collated, analysed and
interpreted.

\section*{Results}
\subsection*{Participant demographics}
There were 20 participants who took part in the first meeting of
the group, and the same number remained throughout the entire
consensus process without dropouts. The demographic profile of the
study participants is shown in Table 1. Our participating healthcare
providers were all experienced in haemophilia care, with 18 of the 20
having 26 years of working in haemophilia care.

\subsection*{Participant response rate and consensus}
The participant response rate in round 2 and round 3 was 100%,
reflecting the commitment of this group to the process. In round 2,
consensus was reached in 33 of the 36 questions (92%).

The top three statements for which there was 100\% expert
agreement are listed in Table 2. The three statements for which there
was no agreement among the study participants are listed in Table 3.
There was general agreement for the remainder of the questionnaire
statements, which are listed in Table 4. The consensus group
identified several potential barriers to home therapy, and the top 10
of these are listed in Table 5.

\section*{Discussion}
Our study participants all agreed that home therapy should target
the paediatric population as a priority over the adult population, that
patients on home therapy should sign an informed consent and an
indemnity, and that patients on home therapy should have round-
the-clock access to a healthcare provider (Table 2).
As indicated in Table 1, a high proportion of participants in the group looked after paediatric patients, which may have skewed the consensus in favour of children. However, there were sufficient non-paediatric participants to have changed this if there was no consensus. The rationale for targeting the paediatric group is the impact home therapy will have in preventing and reducing the haemophilia disease burden in this age group. Owing to poor venous access, many home therapy guidelines do not specifically recommend targeting the paediatric group for home therapy.\(^{[10-11]}\) In a resource-constrained setting such as ours, our group considered the cost-effectiveness of home therapy in terms of volume of CFC to keep at home, as well as the long distances our patients have to travel to reach haemophilia treatment centres. In general, paediatric patients require fewer vials of CFC at home, and more than 10 of the 18 haemophilia treatment centres in SA are servicing patients drawn from rural settings.

The recommendation that patients on home therapy should sign an informed consent and an indemnity came as no surprise. SA is not a litigious society; however, we have seen a steep rise in health-related litigations, particularly in state facilities where patients claim professional negligence.\(^{[10]}\) This recommendation is also in line with increasing societal demands for the practice of ethical medicine in SA and aligns with current hospital practice for patients undergoing procedures to sign indemnity for hospitalisation and consent for all procedures. Once implemented, this recommendation will give patients and healthcare providers the reassurance that home therapy is taken as seriously as any other procedures performed in healthcare facilities.

The above fact links to the third top recommendation, that patients on home therapy should have round-the-clock access to a healthcare provider. Such support is already provided for all haemophilia patients, but becomes mandatory for patients who will be infusing CFC at home, in view of potential complications of poor venous access.

The participant experts in this Delphi process comprised 10 doctors, 9 nurses and a physiotherapist. The equal number of doctor and non-doctor participants in this expert sample reflects the kind of collective decision-making that takes place in the care of patients with haemophilia. While other members of the haemophilia comprehensive care team were considered for inclusion, these panel members were thought to be pivotal in the direct management of haemophilia patients on home therapy. Unlike other Delphi processes in haemophilia, which have often excluded nurses,\(^{[29,32]}\) our process included nurses, as they are critical role players in the instruction of patients on home therapy as well as in their follow-up and treatment monitoring.

Our participating healthcare providers were all experienced in haemophilia care, with 18 of the 20 having 26 years of working in haemophilia care. This is important, as the decisions taken on home therapy have lifelong implications for patients and their caregivers and therefore require considerable experience and knowledge in the haemophilia field. Our Delphi participants did not include administrators or patients, as healthcare providers wanted to reach consensus among themselves before including patients and administrators in the medical decision-making.

The Delphi process participants did not reach consensus on 3 of the 36 statements in the questionnaire. The reasons for failing to reach consensus are fairly clear and reflect the influence of the composition of the group and the age and size of patients they look after. With regard to needle size, the non-consensus ultimately became a disagreement between paediatricians and adult treaters. In the final analysis, this disagreement was an academic exercise, as all CFC comes with pre-packed needles of the same size. On the home environment, paediatricians felt strongly that only knowledgeable, motivated and skilled caregivers should be allowed to participate in the home therapy programme. How much CFC patients take home is ultimately at the discretion of each hospital, and their policy is often informed by whether the patient is on episodic or prophylactic treatment. The generally accepted principle is for patients to have enough CFC to treat at least one bleed at home or to have enough CFC not to miss prophylactic doses until the next scheduled treatment centre visit. Current national and international guidelines are silent on these two aspects.\(^{[29-34]}\)

### Table 1. Demographic profile of the Delphi process participants

<table>
<thead>
<tr>
<th></th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of participants</td>
<td>20</td>
</tr>
<tr>
<td>Participants professions</td>
<td></td>
</tr>
<tr>
<td>Doctor</td>
<td>10</td>
</tr>
<tr>
<td>Nurse</td>
<td>9</td>
</tr>
<tr>
<td>Physiotherapist</td>
<td>1</td>
</tr>
<tr>
<td>Experience in haemophilia care (years)</td>
<td></td>
</tr>
<tr>
<td>≤5</td>
<td>2</td>
</tr>
<tr>
<td>6 - 10</td>
<td>6</td>
</tr>
<tr>
<td>11 - 15</td>
<td>4</td>
</tr>
<tr>
<td>16 - 20</td>
<td>4</td>
</tr>
<tr>
<td>&gt;20</td>
<td>4</td>
</tr>
<tr>
<td>Type of practice</td>
<td></td>
</tr>
<tr>
<td>Adult haemophilia only</td>
<td>3</td>
</tr>
<tr>
<td>Paediatric haemophilia only</td>
<td>7</td>
</tr>
<tr>
<td>Combined adult/paediatric</td>
<td>10</td>
</tr>
<tr>
<td>Number of haemophilia patients in the practice</td>
<td></td>
</tr>
<tr>
<td>20 - 100</td>
<td>12</td>
</tr>
<tr>
<td>101 - 200</td>
<td>2</td>
</tr>
<tr>
<td>201 - 300</td>
<td>1</td>
</tr>
<tr>
<td>301 - 400</td>
<td>1</td>
</tr>
<tr>
<td>401 - 500</td>
<td>2</td>
</tr>
<tr>
<td>&gt;500</td>
<td>2</td>
</tr>
</tbody>
</table>

### Table 2. Top three questionnaire statements for which there was 100% agreement among participants

- Treatment centres should prioritise paediatric patients for home therapy
- Patients on home therapy should sign informed consent and indemnity for participating in the home therapy programme
- Patients on home therapy should be provided with 24-hour round-the-clock support by the treatment centre

### Table 3. Questionnaire statements for which there was no participant consensus

- The home environment most suitable for home therapy
- Sizes of needles supplied to people with haemophilia for home therapy
- Number of doses to be dispensed for home therapy
<table>
<thead>
<tr>
<th>Questionnaire statement</th>
<th>Agreement reached</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Definition of home therapy</td>
<td>• Home therapy is therapy taken by haemophilia patients at home under supervision of the haemophilia care team</td>
</tr>
<tr>
<td>2. Goals and rationale for implementing home therapy need to be defined</td>
<td>• The primary goals of home therapy should be to treat bleeds early or to put patients on prophylaxis in order to reduce bleed rates and associated musculoskeletal complications</td>
</tr>
<tr>
<td>3. Age of starting home therapy</td>
<td>• Home therapy should be started as soon as the caregivers and patients have been instructed on intravenous infusion and deemed competent</td>
</tr>
<tr>
<td>4. Factors to consider before starting home therapy</td>
<td>• Having a fridge to store product at home • Supportive and nurturing home setting • Hygiene conditions at home • Privacy at home • Distance of home from hospital • Ability to contact the treatment team</td>
</tr>
<tr>
<td>5. Patient characteristics that should favour home therapy</td>
<td>• Patients should be competent in IV needle insertion • Patients should be reliable and trustworthy • Patients should be able to identify the complications of IV needle insertion • Patients should understand the importance of a sterile field when inserting the IV needle • Patients should be able to dispose of needles safely • Patients must have passed at least one course on home therapy</td>
</tr>
<tr>
<td>6. Issues of venous access in the home setting</td>
<td>• Central venous and arterial lines should be avoided • A maximum of three unsuccessful venous stabs should require expert intervention</td>
</tr>
<tr>
<td>7. Success indicators for home therapy</td>
<td>• Should include evaluation of annualised bleeding rate • Number of missed injections for patients on prophylaxis • Serial measurement of musculoskeletal haemophilia complications • Number of deaths of patients on home therapy • Improvement in the quality of life of patients</td>
</tr>
<tr>
<td>8. Patients needing particular attention in the home therapy programme</td>
<td>• Poor venous access • Inhibitor patients requiring bypassing agents • Poor home support structure • Non-compliant patients • Patients who have failed home therapy training • Patients unable to handle and store clotting factor concentrate</td>
</tr>
<tr>
<td>9. Important role players required for the support patients on home therapy</td>
<td>• Immediate parent/guardian/spouse • Extended family • Friends and neighbours • Teachers and co-workers • Local clinic sisters • Emergency personnel</td>
</tr>
<tr>
<td>10. Characteristics of haemophilia centres practising home therapy</td>
<td>• Ability to supply unlimited quantities of factor • Availability of staff 24 hours a day to support patients • Adequate treatment centre drug stock level • Ability to monitor and review treatment compliance • Ability to monitor home therapy impact and outcomes • Ability of the treatment centre to train and monitor patients</td>
</tr>
<tr>
<td>11. Characteristics of products used in the home therapy programmes</td>
<td>• The product should be non-thrombotic • The product should be non-immunogenic in previously untreated patients • The product should not be associated with allergy or anaphylaxis • The product should not cause local reaction on skin extravasation • Product overdose should be very well tolerated</td>
</tr>
<tr>
<td>12. Factors to consider when monitoring home therapy</td>
<td>• Volume of product consumption and accountability • Number of bleeds in a patient on prophylaxis • Number of doses required to treat a bleeding episode • Number of times the patient has run out of medication • Number of calls the patient makes to the healthcare providers</td>
</tr>
</tbody>
</table>

IV = intravenous.
There was non-unanimous but significant consensus among the Delphi group members on the remaining statements listed in Table 4. These statements will have the potential for informing policymaking and becoming part of standard operating procedures for all treatment centres in SA. The standardisation of procedures and policies will help reduce the current largely discrepant practices of home therapy healthcare providers.

To our knowledge, this study is the first to examine home therapy closely and to seek consensus among haemophilia healthcare providers in the developing world. The reasons for lack of similar studies in the past remain unclear, but could include the fact that expertise in haemophilia in many resource-constrained settings is limited and the very practice of home therapy might be poorly supported. In contrast, in many developed countries home therapy is regarded as part of the standard of care of haemophilia, and the need for consensus to implement it therefore does not exist.

The recommendations on home therapy would have been incomplete without identifying potential barriers and hurdles to its implementation. The biggest barrier identified from this Delphi process was the healthcare provider being the resource gatekeeper and not allowing patients to self-infuse. We managed to keep our Delphi process free of interference from the healthcare facility administrators, yet there were a number of haemophilia treatment centres in which patients who were adequately trained on home therapy were not allowed to take CFC to infuse themselves at home by either their healthcare team or hospital administrators. It is hoped that raising awareness around this practice will result in rational decision-making and alignment of the dispensing practice with the recommendations of this study.

**Study strengths and limitations**

The study strengths included the cohesiveness of the group, illustrated by the fact that none of the participants were lost during the study. The fact that consensus within the group was reached quickly within two iterations indicated the general convergence of thinking within the group, which it is hoped will translate to uniform policy and practice in SA.

The study had some limitations. The participants were self-selected and therefore may not be representative of haemophilia healthcare providers as a whole in SA. They included only a small number of healthcare providers and did not include patients or administrators, who would have provided a non-healthcare provider perspective to the consensus statements.

**Conclusions**

Through a modified Delphi process, we have been able to answer a number of questions and reached consensus on haemophilia home therapy, including who should be prioritised for home therapy, the prerequisites for home therapy, and barriers to a successful home therapy programme. The adoption of home therapy in haemophilia will result in a number of desirable outcomes that include reduction in bleeding rates, reduction in musculoskeletal complication rates and improvement in quality of life. The consensus statements arising from this Delphi process will serve as a foundation for formulating a home therapy national policy and for narrowing the wide home therapy practice gap currently seen among the haemophilia treatment centres in SA.

**Declaration.** None.

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**Author contributions.** JNM convened the group, formulated the questionnaire, participated in the consensus process, collated responses, wrote the manuscript, and critically reviewed and edited the manuscript. JP conceptualised the idea and obtained support from Novo Nordisk, participated in the consensus process, and critically reviewed and edited the manuscript. YN participated in the consensus process, chaired the feedback meeting, and critically reviewed and edited the manuscript. YG participated in the consensus process, and critically reviewed and edited the manuscript. MV participated in the consensus process, and critically reviewed and edited the manuscript. YN participated in the consensus process, and critically reviewed and edited the manuscript. AI provided methodological support in setting up and running the Delphi process, performed data analysis, and critically reviewed and edited the manuscript.

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**Conflicts of interest.** None.

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Table 5. Potential barriers to home therapy identified by the participants

<table>
<thead>
<tr>
<th>Potential Barriers to Home Therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inadequate budgets in health facilities for home therapy</td>
</tr>
<tr>
<td>Healthcare providers rationing available clotting factor replacement therapy for patients</td>
</tr>
<tr>
<td>Lack of adequate clotting factors in health facilities</td>
</tr>
<tr>
<td>Lack of appropriate protocols and tools to initiate, implement, monitor and evaluate home therapy</td>
</tr>
<tr>
<td>Poor evaluation and monitoring of home therapy outcomes by healthcare providers</td>
</tr>
<tr>
<td>Generally non-compliant patients</td>
</tr>
<tr>
<td>Patients with poor venous access</td>
</tr>
<tr>
<td>Patients who have failed home therapy training</td>
</tr>
<tr>
<td>Patients with poor home support structure</td>
</tr>
<tr>
<td>Patient unable to handle and store clotting factor appropriately</td>
</tr>
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