Studies on the procurement of blood coagulation factor VIII: effects of plasma freezing rate and storage conditions on cryoprecipitate quality

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SUMMARY Plasma was frozen and stored in different ways before processing to cryoprecipitate by a standard thawing technique. Freezing rate was found to be important with slow freezing having a deleterious effect on cryoprecipitate quality. Storage of frozen plasma at constant temperatures for periods up to six months had no effect on the quality of cryoprecipitate, with no difference being found for plasma stored at −20°C or −40°C. Subjecting frozen plasma to deliberate temperature fluctuations resulted in a considerable increase in the amount of fibrinogen recovered in cryoprecipitate, with the factor VIII yield being relatively unaffected.

Blood bank production of cryoprecipitate still constitutes an important source of therapeutic material for treatment of haemophilia, while bulk cryoprecipitation is the preliminary step in 98% of the world’s production of factor VIII concentrate. Any factors influencing the quality of cryoprecipitation thus have an important bearing on the logistics of haemophilia care. Several studies have established that fast plasma thawing in a low temperature waterbath results in higher factor VIII yields than the original overnight thaw in air. The introduction by Mason of a modified form of fast plasma thawing, in which thawed supernatant plasma was siphoned over leaving the cryoprecipitate in the original plastic pack, further optimised this aspect of cryoprecipitate production. This method results in factor VIII yields of 70% compared with 40% using other techniques, but it is labour intensive and considered by some to be too cumbersome to allow routine production.

Although much data has been published regarding thawing methods, uncertainty exists about the optimal way of freezing plasma. Placement of fresh plasma packs in a −30°C or −20°C freezer has been reported to lead to decreased factor VIII yields in cryoprecipitate compared with plasma frozen faster in blast freezers. Placement in −40°C freezers has been reported to give yields similar to faster freezing techniques such as −80°C blast freezers or placement in liquid nitrogen.

Storage of frozen plasma before processing to cryoprecipitate is another aspect that has received little attention. Although early studies suggested that plasma factor VIII clotting activity (VIII:C) is labile at −20°C, subsequent work, using more reliable assays, reported no loss in VIII:C for periods of up to 20 months at −20°C. As far as production of cryoprecipitate goes, however, VIII:C is not the only parameter of importance; changes in other proteins have been reported to occur during frozen plasma storage and to result in processing difficulties during subsequent production of cryoprecipitate and factor VIII concentrate.

In this study a modification of the thaw-siphon technique has been used to study the effects of plasma freezing rate and storage conditions on cryoprecipitate yield and purity.

Material and methods

Plasma used in this study was prepared from normal citrate phosphate dextrose blood donations within 3 h of donation. Plasma pools of six donations were used in each experiment to avoid the variations in VIII:C content found in different individuals. The plasma was pooled in a 21 pack and aliquots of 200 ml were distributed into 300 ml Fenwall R2011 transfer packs. The plasma was then frozen and stored as described below in one of two ways. Either all packs were subjected to fast freezing and then

Accepted for publication 12 December 1984

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aliquots stored at −20°C or −40°C and processed to
cryoprecipitate at 16 h, three months, or six months;
or three packs from each pool were used to study the
effect of freezing rate. These packs were further
used to study the effect of temperature insult during
storage (fast frozen units) and time of frozen storage
(slow frozen units). For each comparison six distinct
plasma pools were used. Statistical analysis was
based only on results obtained from such split pools
processed as follows.

SLOW FREEZING
Packs of plasma were placed in thin aluminium cassettes
and placed in a −40°C cabinet freezer. The
cassettes were placed vertically and the packs had
the outlet ports down.

FAST FREEZING
Packs were placed in cassettes as above but were
frozen in a −70°C ethanol bath which had been
cooled using liquid nitrogen or solid carbon dioxide.

FREEZING RATES
For both these freezing techniques, plasma tempera-
ture was monitored with a thermocouple (Comark
Instruments) placed in the middle of a pack, which
was connected to a pen recorder (Vitatron).

STORAGE STUDIES
Plasma pools prepared as above were frozen in
−70°C ethanol or −40°C air and stored at different
temperatures and for different time periods as
described in the Results section. Daily checks
ensured that the storage temperatures were within
reasonable limits (±4°C) of the temperatures stated.
In studies involving temperature insult plasma packs
frozen to −70°C were subjected to deliberate
fluctuations in temperature by placing at +4°C until
the core temperature reached −5°C.

CRYOPRECIPITATE PREPARATION
This was as described by Mason et al16 with some
modifications. Instead of the tensioning device
described by Mason et al, elastic bands were
stretched between metallic rods placed through the
slits in the packs’ lateral edges so as to maintain the
shape of the pack during thawing. On completion of
thawing, the cryoprecipitate was dissolved at 37°C in
its own residual supernatant plasma and samples of
cryoprecipitate and cryosupernatant were stored at
−40°C for subsequent assay.

Factor VIII:C was assayed using a one stage clot-
ting assay,17 factor VIII coagulant antigen (VIII:C
CAg) was assayed immunoradiometrically,18 and
factor VIII related antigen (VIIIIR:Ag) was deter-
mined by electroimmunoassay,17 except for samples
of cryosupernatant plasma, which were assayed
immunoradiometrically.19 Fibrinogen was assayed
by electrophoresis,17 and total protein by the
Biuret method.20

Results

Fig. 1 shows the temperature profiles of plasma
packs frozen under the two different sets of condi-
tions studied. The average cooling rates were
−5°C/minute and −0.33°C/minute for fast and slow
frozen plasma respectively. Plasma frozen under
these conditions was processed to cryoprecipitate
after overnight storage at −40°C. It was immedi-
ately noticeable that plasma frozen slowly pro-
ced a steady flux of particulate material in the siphon
lines during thawing. On the other hand, fast frozen
plasma produced a clear supernatant with no par-
ticles, unless thawing was allowed to proceed until
the residual cryoprecipitate also siphoned over. Fig. 2
shows that slow freezing resulted in a slightly lower
recovery of total VIII:C (expressed as the sum of the
VIII:C recovered in cryoprecipitate and cryo-
supernatant) on thawing the plasma; this difference
was not significant (p > 0.1). In slowly frozen plasma,
however, only 47% of the residual VIII:C was re-
covered in the cryoprecipitate after thawing. This
poor cryoprecipitability of factor VIII in slowly
frozen plasma was confirmed by assays of other factor
VIII related activities in cryoprecipitate and
cryosupernatant (Fig. 2). Fast frozen plasma gave
much better results: 425 units of VIII:C/kg plasma
was recovered in cryoprecipitate compared with 318
units/kg for slowly frozen plasma (p < 0.05). Plasma
frozen fast also gave yields of fibrinogen in cryo-
precipitate of 824 mg/kg plasma, compared with 522

\[
\begin{align*}
\text{Temperature recorded in plasma packs frozen in different media.}
\end{align*}
\]
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Fig. 2 Distribution of factor VIII related activities in plasma fractions derived from plasma frozen at different rates. Mean ± SD of six experiments expressed as U/kg of starting plasma.

mg/kg for slowly frozen plasma (p = 0.01). The appearance of particulate material during thawing, as well as the higher levels of factor VIII related activities in the cryosupernatant derived from slowly frozen plasma, suggests that cryoprecipitate from such plasma is not efficiently retained during the thawing process and is lost in the supernatant during siphoning.

The Table summarises data on plasma frozen in different ways and stored under constant temperatures for different periods. Data for fast frozen plasma in this series of experiments are based on distinct plasma pools from those used for the freezing rate experiments shown in Fig. 2. As long as constant temperature was maintained, there was no significant difference in cryoprecipitate factor VIII and fibrinogen for plasma stored at -20°C and -40°C; storage of plasma for up to six months at either temperature also had no deleterious effect. While there is some suggestion of reduced yields in cryoprecipitate prepared from plasma stored frozen for six months, it would be necessary to compare a large number of plasma pools to establish if this is significant.

Fig. 3 shows the temperature fluctuations that were recorded in the core of a plasma pack sub-

<table>
<thead>
<tr>
<th>Freezing rate</th>
<th>Period of storage</th>
<th>Temperature of storage</th>
<th>Total VIII:C recovered (precipitate + supernatant) (U/kg plasma)</th>
<th>VIII:C precipitate yield (U/kg plasma)</th>
<th>Fibrinogen precipitate yield (mg/kg plasma)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fast*</td>
<td>16 h</td>
<td>-20°C</td>
<td>623 ± 108</td>
<td>426 ± 80</td>
<td>605 ± 172</td>
</tr>
<tr>
<td></td>
<td>3 mo</td>
<td>-20°C</td>
<td>698 ± 123</td>
<td>500 ± 118</td>
<td>609 ± 187</td>
</tr>
<tr>
<td></td>
<td>6 mo</td>
<td>-20°C</td>
<td>690 ± 82</td>
<td>416 ± 67</td>
<td>538 ± 115</td>
</tr>
<tr>
<td></td>
<td>3 mo</td>
<td>-40°C</td>
<td>698 ± 125</td>
<td>493 ± 116</td>
<td>607 ± 292</td>
</tr>
<tr>
<td></td>
<td>6 mo</td>
<td>-40°C</td>
<td>718 ± 102</td>
<td>522 ± 77</td>
<td>577 ± 51</td>
</tr>
<tr>
<td>Slow†</td>
<td>16 h</td>
<td>-40°C</td>
<td>730 ± 101</td>
<td>449 ± 47</td>
<td>542 ± 47</td>
</tr>
<tr>
<td></td>
<td>3 mo</td>
<td>-40°C</td>
<td>610 ± 101</td>
<td>318 ± 111</td>
<td>522 ± 270</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>611 ± 183</td>
<td>306 ± 62</td>
<td>502 ± 79</td>
</tr>
</tbody>
</table>

* -70°C liquid N₂/ethanol.
† -40°C deep freeze.
‡ Achieved by placement in deep freezes of the appropriate temperature. Results show mean ± SD for six different experiments, each using a pool of six donations, aliquots of which were treated in the different ways described. Distinct sets of pools were used for the fast freezing and slow freezing experiments and these results are thus not directly comparable. Fig. 2 shows the results of studies designed to test the effect of freezing rate.
method described by Mason results in considerable improvements in cryoprecipitate factor VIII yield. In the modified method described subsequently, and used in this study, some of this increased yield is sacrificed to obtain improved purity. Thus, while a specific activity of 0-09 U/mg total protein is achieved using conventional thaw-siphoning, the modified technique gives up to 0-5 U/mg. In the present study, a specific activity of 0-19 U/mg was achieved, which is lower than found by Mason et al., possibly as a result of less efficient tensioning leading to retention of excess plasma protein with the cryoprecipitate. As has been pointed out previously, the use of split plasma pools in comparing different modes of cryoprecipitate production is essential. While differences in the assay and processing of various pools are, to some extent, unavoidable, these were minimised in the present study by using well characterised established techniques.

Previous studies using the thaw-siphon technique all used plasma frozen by fast freezing methods. The importance of fast freezing is confirmed by the present study. Placement of plasma in a -40°C freezer, a method found to be adequate for plasma processed by conventional fast freezing in liquid nitrogen/ethanol to -40°C and held at -40°C for seven days (O). Insulted once by warming to -5°C on day 2 and then replaced in -40°C storage (▲). Insulted twice after two and five days (○).

Fig. 3 Temperature recorded in a plasma pack subjected to temperature insult. Control: frozen in liquid nitrogen/ethanol and held at -40°C for seven days (O). Insulted once by warming to -5°C on day 2 and then replaced in -40°C storage (▲). Insulted twice after two and five days (○).

Discussion

In this study a standard thaw-siphon method was used to produce cryoprecipitate. The original

Fig. 4 Effect of temperature insult (warming and refreezing) on cryoprecipitate preparation. Mean and SD for six experiments. All units were processed to cryoprecipitate seven days after initial plasma freezing. (a) Yield of VIII:C in plasma fractions. (b) Yield of fibrinogen in cryoprecipitate. (c) Specific activity of VIII:C (U/mg fibrinogen) in cryoprecipitate.
thawing,\(^1\) \(^1\)\(^2\) resulted in decreased yields of factor VIII related activities and of fibrinogen in cryoprecipitate. Particulate matter was observed as soon as slowly frozen plasma started thawing, and this phenomenon was accompanied by a loss of factor VIII related activities to the supernatant. This suggests that cryoprecipitate formation and harvesting depends on the mode of freezing. Fast freezing is necessary for optimal yields of factor VIII, at least by the technique used here.

Within the limits tested, the storage time or temperature of frozen plasma did not affect cryoprecipitate quality, so long as steady storage conditions were maintained. The finding that –20°C is adequate is important as storage at this temperature is much less expensive than at colder temperatures. Although it has been suggested that storage below the eutectic point is to be preferred,\(^2\)\(^1\)\(^1\)\(^1\) no evidence for this was found in the time scale of the present study. Deliberate temperature insult during storage, however, had a pronounced effect on cryoprecipitate quality. Although VIII:C yields in cryoprecipitate from insulin plasma were not severely affected, the total amount of VIII:C recoverable declined by 20% owing to a sharp drop in the VIII:C recovered in cryosupernatant. The increased lability of cryosupernatant VIII:C compared with cryoprecipitate VIII:C has been documented\(^2\)\(^2\)\(^2\) and it seems that the deleterious effects of temperature insult result in loss of cryosupernatant VIII:C activity. The striking increase in cryoprecipitate fibrinogen that results from temperature insult may be one of the reasons for the variation in cryoprecipitate mass that has been noted during plasma fractionation.\(^3\)\(^3\)\(^3\)\(^3\)\(^3\)

Temperature fluctuations such as those deliberately engineered in the present study may occur during storage and transportation of bulk frozen plasma, although warming of large plasma blocks is bound to be less rapid than for the smaller blocks used in the present study. Increased cryoprecipitate fibrinogen is responsible for difficulties in extracting and further processing factor VIII concentrate and may result in poor filterability and solubility. The storage temperature profile is thus an important variable in determining the quality of plasma destined for fractionation to factor VIII products.

AF was a recipient of a British Council Grant (MLT/781/01) during the course of this study and was on study leave from the Blood Transfusion Unit, St Luke’s Hospital, Malta.

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doi: 10.1136/jcp.38.4.433

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