Cryopreservation of *Mycobacterium tuberculosis* Complex Cells

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Abstract

Successful long-term preservation of *Mycobacterium tuberculosis* (MTB) cells is important for sample transport, research, biobanking, and the development of new drugs, vaccines, biomarkers, and diagnostics. In this report, *M. bovis* Bacillus Calmette-Guérin (BCG) and *M. tuberculosis* H37Ra were used as models of MTB complex strains to study cryopreservation of MTB complex cells in diverse sample matrices at different cooling rates. Cells were cryopreserved in diverse sample matrices, namely phosphate buffer saline (PBS), Middlebrook 7H9 medium with or without added glycerol, and human sputum. Efficacy of cryopreservation was quantified by microbiological culture and microscopy with BacLight live/dead staining. In all sample matrices examined, the microbiological culture results showed that cooling rate was the most critical factor influencing cell viability. Slow cooling (a few degrees Celsius per minute) resulted in much greater MTB complex recovery rates than rapid cooling (direct immersion into liquid nitrogen) (*p*<0.05). Among the three defined cryopreservation media (PBS, 7H9, and 7H9+glycerol), there was no significant differential effect on viability (*p*=0.06~0.87). Preincubation of thawed MTB complex cells in 7H9 broth for 20 hours before culture on solid Middlebrook 7H10 plates did not help the recovery of the cells from cryoinjury (*p*=0.14~0.71). The BacLight live/dead staining kit, based on SYTO9 and propidium iodide (PI), was also applied to assess cell envelope integrity after cryopreservation. Using the kit, similar percentages of “live” cells with intact envelopes were observed for samples cryopreserved under different conditions, which was inconsistent with the microbiological culture results. This implies that suboptimal cryopreservation might not cause severe damage to cell wall and/or membrane, but instead cause intracellular injury, which leads to the loss of cell viability.

**Keywords:** cryopreservation, *Mycobacterium tuberculosis* (MTB) complex, Bacillus Calmette-Guérin (BCG), *M. tuberculosis* H37Ra, cooling rate, preincubation
Introduction

Tuberculosis (TB) is the second leading cause of death from an infectious disease worldwide (after HIV/AIDS). Millions of people die of this disease every year (1.2-1.5 million deaths in 2010). An active TB patient can infect 10-15 persons in a year without restriction or treatment.

TB prevention and control, including diagnosis, treatment and drug development is one of the major tasks of World Health Organization (WHO) and many governments, especially in developing countries. (24)

Timely and correct diagnosis of TB is still a challenge. Often collected samples must be shipped to centers with clinical laboratory facilities, well-trained personnel and specialized working environments for biosafety. In this context, cryopreservation can be a useful option. Biobanking of Mycobacterium tuberculosis (MTB) is also important in fundamental biological research and for the development of new drugs, vaccines, biomarkers and diagnostic tests. Clinical and laboratory samples are often exchanged among institutions for comparative multicenter studies. Therefore quality control of transported samples is a very important issue.

There have been few studies on the optimization of MTB cryopreservation. Clinical MTB samples such as sputum are stored at room temperature, but only for a few days (4, 10, 20). Viability of some bacteria, such as MTB and M. bovis, can be severely affected during shipping by storage conditions (14). Papers published a few decades ago demonstrated long-term preservation of MTB samples at -70 °C, but little information on optimization of the cryoprotective medium and freezing procedure was provided (9, 11-14). Therefore, optimization of MTB cryopreservation remains an unfilled need, especially for situations requiring quantification of viable MTB cells in samples with low cell density.

Since M. bovis Bacillus Calmette-Guérin (BCG), a non-virulent strain of the M. tuberculosis complex, is very close to M. tuberculosis in terms of cell physiology, it was used as the model to
optimize the cryopreservation protocol in the first part of this study. Common cryopreservation media, effect of a cryoprotective agent (glycerol) and cooling rate were studied. It has been suggested that transient incubation of cells in broth medium after exposure to stress environments, such as heating, freezing/thawing or ultraviolet (UV) light, helps to repair the injury for some cell types (1, 8, 15, 23). Therefore, the effect of transient incubation (preincubation) on cryopreserved BCG cells was also explored. To conclude the study, the optimized protocol was applied to M. tuberculosis H37Ra cells spiked into a natural, undefined sample matrix, namely human sputum.

Materials and Methods

BCG cell preparation and concentration assessment

BCG cells were cultivated to late log/early stationary phase in 15 ml conical tubes containing 5 mL Difco Middlebrook 7H9 broth with 10% ADC enrichment and 0.05% Tween 80 (BD Diagnostics, Sparks, MD) (henceforth referred to as 7H9) at 37 °C without agitation for 3-4 weeks. Cells were harvested by centrifugation for 5 min at 5000rpm and then resuspended in an equal volume of fresh 7H9. Cell concentration was assessed by optical density measurement at 580 nm with a spectrophotometer (Molecular Devices Spectramax Plus, Sunnyvale, CA). According to previously published growth measurements an A580 of 0.1 units is equivalent to 6.3x10⁷ CFU per mL Mycobacterium cells (3).

BCG cryopreservation

BCG cells were cryopreserved in three different media: Phosphate-buffered saline (PBS), 7H9, and 7H9 + glycerol (EMD Chemicals, INC., Gibbstown, NJ). For PBS and 7H9, BCG cells were diluted in PBS or 7H9 respectively to an estimated cell density of 1e8 CFU/mL (based on optical density). For 7H9 + glycerol, BCG cells were first diluted with 7H9 to 5e8 CFU/mL, and then 7H9 + glycerol (pre-cooled to 4 °C) was slowly added dropwise to final volume ratio of 4:1. The
The final glycerol concentration in the freezing sample was 4% [v/v]. The cell suspension was agitated gently for osmotic equilibration during medium addition. After medium addition, the cell suspension was stored at room temperature for 15 minutes for equilibration before cryopreservation. For all three groups, the final cell concentration was 1e8 CFU/mL. This high cell density is typical of many cryopreserved samples and it enabled Live/Dead staining and microscopic analysis. Cryogenic freezing vials (2 mL, Fisher Scientific, Pittsburgh, PA) were used for cryopreservation, each containing 0.5 mL of the cell suspension.

For comparison between different cooling rates, two freezing methods were implemented. One was slow cooling in a “Box-in-Box” (BIB) system that was developed in our group, and the other was rapid cooling by direct immersion of samples into liquid nitrogen (LN2). In the BIB system (Fig 1), a thermal insulation material made of polyethylene fills an aluminum enclosure. The thickness of the polyethylene layer was designed for a low cooling rate and then demonstrated empirically (21). For cryopreservation in BIB, vials were placed in the slots cut into the polyethylene foam. The system was then closed and placed into a -80 °C freezer overnight. Dummy samples with only cryoprotective medium were treated the same way. T-type thermocouples (SA-1T; Omega, Stamford, CT) were inserted inside the dummy samples to record temperature history during cooling. After equilibrium in the -80 °C freezer overnight, the samples were transferred into liquid nitrogen (LN2). The protocol for fast cooling was to immerse the sample vials into LN2 directly, and record temperature history via dummy samples.

**Thawing of cryopreserved BCG cells**

For sample thawing, the frozen samples were removed from LN2 and quickly immersed into a 37 °C water bath. They were agitated gently until no crystals remained. Then the cells were assessed for viability by microbiological culture and live/dead staining as below.

**Microbiological culturing, with and without preincubation**
The thawed cells were diluted with 7H9 broth to a concentration of 4000 CFU/mL. Before culturing, the cells were divided into two groups. One group was plated on 7H10 agar directly (without preincubation), and the other was pre-incubated in 7H9 at 37 °C with 5% CO2 and 100% humidity for 20 hours before plating in order to study the preincubation effect. For plate culturing, Difco Middlebrook 7H10 agar with 10% (v/v) OADC enrichment (BD Diagnostics, Sparks, MD) (7H10) was used. Cell suspension of 250 µL with a predicted concentration of 4000 CFU/mL was spread on each plate. Plates were sealed in plastic bags (BD GasPak EZ; BD Diagnostics, Sparks, MD) and incubated at 37 °C with 5% CO2, and 100% humidity. After incubation for about 24 hours, the petri dishes were inverted in order to prevent agar from desiccation during extended culture.

These experiments were repeated three times with three separate batches of BCG cells. CFUs were counted at several time points post-inoculation.

Live/Dead staining assessment

A BacLight Live/Dead staining kit (Invitrogen, Grand Island, NY) was used to assess cell envelope integrity of cryopreserved cells. The staining was performed as specified by the user manual, briefly: 1.5 µL live/dead staining reagent was added to 0.5 mL cell suspension, which was vortexed, incubated at room temperature for 15 minutes, and then observed under fluorescence microscopy (Nikon Instruments INC, Melville, NY).

Cryopreservation of M. tuberculosis H37Ra cells in sputum

The experimental design of H37Ra cryopreservation in sputum is illustrated in Fig. 2. H37Ra cells were spiked into sputum samples that had been collected for diagnostic purposes from anonymous human donors with no tuberculosis exposure (Bioreclamation Inc, Hicksville, NY). Estimated final cell concentrations were 1e7 CFU/mL. Then the samples in cryogenic freezing vials (0.5 mL in each vial) were frozen with three different methods: BIB in a -80 °C freezer, in...
liquid nitrogen (LN2) directly, or in a -80 °C freezer directly. After fast-thawing in a 37 °C water bath, samples were treated with N-acetyl-L-cysteine and sodium hydroxide (NALC-NaOH) following the standard digestion-decontamination procedures for TB specimen processing (6). The samples were centrifuged at 3000g for 20 min, and the pellets were resuspended in 7H9 for microbiological culturing as described above. The experiments were repeated twice.

In order to differentiate the effects of NALC-NaOH treatment and cryopreservation, three control experiments were conducted, as shown in Fig 2. In Control 1, H37Ra cells were spiked into 7H9 and then cultured (without cryopreservation, NALC-NaOH treatment, or centrifugation). In Control 2, H37Ra cells were spiked into 7H9, and the samples were treated with PBS buffer following the procedure of NALC-NaOH treatment (with PBS in place of NALC-NaOH), centrifuged, and then cultured (without cryopreservation and chemical influence of NALC-NaOH). The difference between Control 1 and Control 2 is the effect of centrifugation on cell loss. In Control 3, H37Ra cells were spiked into sputum, and the samples were treated with NALC-NaOH, centrifuged, and then cultured without cryopreservation. Comparison between Control 2 and Control 3 indicates the effect of NALC-NaOH treatment.

Statistical analysis

The results are presented as average ± standard deviations. The statistical analysis was performed with a two-way ANOVA test for the comparison between different conditions. The level of significance was set as p<0.05.

Results

Temperature record and cooling rate during freezing

Typical temperature profiles in the cooling process were recorded for dummy samples in the BIB system, in liquid nitrogen (LN2), and in a -80 °C freezer directly, and the cooling rates were
computed. In the BIB system, a slow cooling rate of -1 to -3.5 °C/min was observed. When
samples were immersed into LN2 directly, the cooling rate ranged between -100 and -
400 °C/min. For samples in a -80 °C freezer directly, the cooling rate ranged between -10 and -
20 °C/min. For freezing directly in the freezer, the cooling rate was variable and depended on
the location of the samples and heat transfer situation inside the freezer.

Viability of BCG cells after freezing and thawing

CFU numbers were counted at various time points after plating. After incubating for two weeks,
colonies started emerging, while after four weeks the CFU numbers reached stability. There
was little or no additional colony formation between 4 and 8 weeks culture.

The CFU recovery rates compared to control samples are shown in Fig 3. (In the first
experiment cells were cultured for three weeks and in the second and third experiments for four
weeks). The results indicate that the cooling rate is one of the most critical factors for BCG cells
to survive freezing and thawing. A slow cooling rate enabled about one order of magnitude
greater CFU recovery than a fast cooling rate (p<0.05) in all freezing media tested. There was
no significant difference between any of the three freezing media (p=0.06~0.87).

CFU counts of cryopreserved BCG with and without preincubation before plating are also shown
in Fig 3. For cryopreserved BCG cells, preincubation did not significantly enhance the cells’
recovery (p=0.14~0.71).

In conclusion, among the experimental conditions examined, only cooling rate exhibited a
significant and consistent effect on BCG cryopreservation.

Live/Dead staining of BCG cells

The BacLight Live/Dead staining kit, which differentially stains bacteria with intact vs.
compromised cell envelopes, was used to assess cell envelope integrity of cryopreserved cells.
For all treatment groups, the percentage of intact cells after cryopreservation and thawing was evaluated with the BacLight live/dead staining kit (Fig 4). Among different freezing media and cooling rates, there was no significant difference in the percentage of intact (“live”) cells (p=0.14~0.99). Compared to controls (fresh cells without cryopreservation), percentages of intact cells appeared high under all conditions, despite the sharply reduced viability (assessed by culture) of cells that were rapidly frozen in LN2.

The live/dead staining kit indirectly assesses viability in terms of cell envelope integrity. To validate the kit's performance on BCG cells, it was also applied to mixtures of fresh and heat-treated BCG cells and compared with the results of microbiological culturing. BCG cells were heated at 80 °C for 35 min (inactivated cells), and then the treated cell suspension was mixed with fresh (viable) cells. The two cell suspensions had the same cell concentrations (estimated by optical density) and were mixed with varying volume ratios of 0:5, 1:4, 2:3, 3:2, 4:1 and 5:0. The mixtures were divided into two groups. One was assessed by BacLight Live/Dead staining, and the other one was cultured on 7H10 agar. For microbiological culturing, each plate received 250 µL of cell suspension with a predicted starting concentration of 4000 CFU/mL (i.e. 1000 cells/plate, including both fresh and heat-treated). The CFU numbers were counted after culture for 28 days. As shown in Fig. 5, live/dead staining yielded results that closely correlated with microbiological culturing results for heat-treated BCG cells. In contrast, a similar experiment of live/dead staining for the mixtures of fresh and cryo-killed BCG cells (fast cooled in LN2) was also conducted. However, the results did not show significant difference among the mixtures of fresh and cryo-killed cells with different volume ratios (0:5, 1:4, 2:3, 3:2, 4:1 and 5:0) (Fig.5).

Viability of M. tuberculosis H37Ra cells after cryopreservation in sputum

The CFU counting results of cryopreserved H37Ra cells in sputum are presented in Fig. 6. The results show that treatment with NALC-NaOH, centrifugation, and freeze-thaw processing can
result in viable cell loss. Comparison between Controls 1, 2 and 3, indicates that treatment with NALC-NaOH and centrifugation can each cause viable cell loss by about one decade. As the results of BCG cryopreservation in defined matrices, cooling rate was a critical factor for MTB cryopreservation in sputum. Fast cooling in LN2 reduced viability of MTB cells to a greater extent than slow cooling in BIB (p<0.05).

Direct freezing of MTB samples in a -80 °C freezer has been used in clinics and research due to its low cost and convenience. MTB cell recovery rate after cryopreservation was compared between direct freezing in BIB, and direct LN2 immersion. Of these conditions, the higher survival was observed when BIB was used (p<0.05). However, the difference between freezing in BIB and directly in the -80 °C freezer was significant in the 1st experiment (p<0.05), but not in the 2nd experiment (p=0.73). This might be due to the fact that freezing rates of samples in a -80 °C freezer vary significantly with location inside the freezer, freezer operation, and heat transfer conditions. The BIB system can provide a more reproducible slow-cooling environment for cryopreservation.

**Discussion**

Currently there is no explicitly standardized protocol for the cryopreservation of clinical TB samples. Samples are often placed directly into a -80 °C freezer. However, this practice leads to variable cooling rates dependent on many factors including the sample container, volume of samples, and heat transfer condition inside the freezer. Also the cooling rate in the freezer is not optimally slow for MTB cryopreservation, as shown in the results. In this work, we employed a simple passive cooling device—“Box-in-Box”, which enables a consistent and reproducible slow cooling rate. The BIB method is inexpensive, easy to operate, reliable, and delivers a reproducible cooling rate. The BIB device has been successfully used for cryopreservation of stem cells (21, 25, 26). The results in this paper showed that it delivers the highest recovery of MTB after cryopreservation.
In cryopreservation, according to the “two factors” hypothesis, cryoinjury to cells during freezing consists of “ice injury” and “solution injury” (16, 17). Many parameters might influence the cell recovery rate after freezing and thawing, e.g. cooling rate, the presence of cryoprotective agent (CPA), CPA addition and removal, and thawing process. The optimum procedure also depends on cell properties, such as the cell membrane permeability to water and CPA, the ratio of cell membrane area to cell volume (A/V), and osmotically inactive intracellular volume fraction \(V_b\).

The results of this work showed that slow cooling is much better than fast cooling for cryopreservation of MTB cells. This may be explained by the properties of MTB cells. The *Mycobacterium* cell envelope has low permeability to most compounds including water (“permeability barrier”), because its outer leaflet is composed of giant saturated fatty acids with extremely low fluidity (18, 19). During freezing, extracellular water is frozen first, which causes the osmolality of extracellular solution to increase. Due to the unbalance of water chemical potential between the intra- and extracellular environments, intracellular water transports across the cell membrane and wall to the outside. This increases the osmolality of intracellular solution, decreasing its freezing point, and hindering intracellular ice formation (IIF). For MTB cells, water may permeate across the cell envelope more slowly than the envelopes of other cell types. More time would be required to reach equilibrium between the inside and outside of the cells during freezing. Slow cooling might mitigate the intracellular ice injury in MTB cryopreservation.

Further research on MTB cell membrane and wall properties, including quantitative cell membrane permeability to water, would be helpful for cooling protocol optimization and interpretation.

The cryopreservation medium and specific CPAs are additional important factors that may influence the cell recovery. Generally CPAs such as glycerol are added before freezing to reduce intracellular ice formation. The standard formulation of 7H9 broth contains glycerol, but at a very low concentration (0.2%, v/v). Due to its widespread use in cryopreservation, glycerol
was evaluated as the CPA in this study. However, the results revealed no significant benefits of glycerol (p=0.11~0.87). In the future, additional CPAs (such as dimethyl sulfoxide) should be tested.

Compared to PBS, 7H9 broth is a complex medium with differences in composition, pH, and osmolality. It contains many kinds of electrolytes, a low concentration of glycerol, and nutrients with pH=6.8 and osmolality of about 168mOsm/Kg H₂O. In contrast, PBS is pH=7.2~7.4, and its osmolality is about 290mOsm/Kg H₂O. The osmolality was measured by vapor pressure osmometer (Wescor INC., Logan, Utah). Despite these differences, we did not observe a significant difference between BCG cells cryopreserved in 7H9 and PBS (p=0.11~0.84).

Compared to cooling rate, cryopreservation medium and CPA had limited impact. The results indicated that for convenience and simplicity, MTB cells could be cryopreserved well in 7H9, PBS, or possibly other media, as long as the cooling rate is slow (generally recovery rates >50%). This eliminates the steps of CPA addition and removal before and after cryopreservation, which add complexity and increase the risk of sample contamination.

When cells are exposed to stressful environments, e.g., cooling, heating, radiation, or desiccation, cellular functions may be impaired. It has been observed that for some cells, preincubation in broth or liquid medium for some time after exposure to the stress may help repair the injuries (1, 8, 15, 23), however others doubt its effectiveness (7). In the present work, preincubation did not improve the recovery of cryopreserved BCG cells from cryoinjury.

The mechanism of cryoinjury during rapid cooling did not appear to involve severe damage to the cell envelope. By live/dead staining, rapidly cooled cell suspensions with low viability did not significantly differ from slowly cooled suspensions with high viability. The BacLight live/dead staining kit contains SYTO 9 and propidium iodide (PI). PI can penetrate only cells with compromised cell envelope integrity, while SYTO 9 can penetrate both intact cells and cells with
compromised cell envelopes. Thus, intact cells emit green fluorescence while compromised cells emit red fluorescence under fluorescence microscopy, i.e., the live/dead staining kit indirectly assesses viability in terms of cell envelope integrity. After suboptimal (rapid) freezing and thawing, many BCG cells might retain intact walls and membranes, while sustaining enough intracellular injury to compromise their ability to grow. This contrasts with BCG cells heated at 80 °C for 35 minutes. For these heat-killed cells, live/dead staining and microbiological culture results correlated with each other, consistent with cellular inactivation mechanisms that include cell envelope damage. The commercially available BacLight live/dead staining kit was designed primarily for use on bacteria, but lacks enough information on efficacy when applied to MTB cells. It was previously reported that the staining pattern of the kit depends on the bacterium type, and even the way of treatment to the bacteria (such as heating, UV exposure), especially for cells in intermediate states between viable and inactivated states (2, 5, 22). Additional evaluation for the kit should be conducted for MTB viability assessment.

Conclusions

Among the variables tested, cooling rate was the most critical factor for the cryopreservation of Mycobacterium tuberculosis complex cells. MTB complex cells were well cryopreserved in PBS buffer, Middlebrook 7H9 with and without added glycerol, or sputum if a slow cooling rate was applied. Preincubation of frozen/thawed BCG cells in 7H9 broth before culturing on solid 7H10 agar did not help the cells repair cryoinjury. Cell inactivation by fast cooling was not associated with a compromised cell envelope, as indicated by the results of microbiological culture and live/dead staining.

Acknowledgements

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References


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Fig 1 Box-in-Box (BiB) system for slow cooling. (A) An opened BiB, in which sample vials are placed in one slot. (B) Cross-section of the BiB system.
Cryopreservation of H37Ra in sputum

Spike H37Ra into sputum

Freeze in BIB

Freeze in LN2

Freeze in -80°C freezer

Thaw

Thaw

Thaw

Treat with NALC-NaOH

Treat with PBS (similar to NALC-NaOH treatment)

Centrifuge, resuspend

Centrifuge, resuspend

Centrifuge, resuspend

Centrifuge, resuspend

Centrifuge, resuspend

Culture

Culture

Culture

Culture

Culture

Culture

Fig 2 Experiment design of H37Ra cryopreservation in sputum
Fig 3 Microbiological culture results of cryopreserved and thawed BCG cells.
Fig 4 Intact BCG cell percentages after cryopreservation assessed by live/dead staining (mean±STD, n=14). Control: fresh cells.
Fig 5 Correlation between live/dead staining and microbiological culturing results when applied to heat-treated (at 80 °C for 35 min) and cryo-killed (fast cooled in liquid nitrogen) BCG cells. (▲) CFU numbers for mixtures of fresh and heat-killed BCG cells; (■) Intact cell percentage for mixtures of fresh and heat-treated BCG cells; (□) Intact cell percentage for mixtures of fresh and cryo-treated BCG cells.
Fig 6 CFU results of cryopreserved and treated H37Ra cells. (A) CFU recovery rates (%) after cryopreservation, NALC-NaOH treatment and centrifugation, normalized by control 1. (B) Influence of cooling rate on CFU recovery rates (%) after cryopreservation, normalized by control 3. (n=6)