Pathogen inactivation: prospects and pitfalls

**Keywords:** HIV, HBV, HCV, blood transfusion, pathogen inactivation, UVA

**Introduction**

While the spread of viral diseases like HIV, HBV, HCV via blood transfusion have been controlled to a large extent but the threats from emerging pathogens and the bacterial contamination of platelets concentrates (PC) remains a major threat with serious clinical consequences. In contrast to the well-established pathogen inactivation strategies for fresh frozen plasma using the solvent-detergent procedure or methylene blue and visible light, the bench-to-bedsite translation of novel pathogen inactivation technologies for cell-containing blood components such as platelets and red blood cells are still underway.¹

Pathogen inactivation (PI) systems have been developed that have proven to be effective against numerous bacteria, viruses, and parasites. Two main systems have emerged to treat platelet concentrates: treatment with psoralen ultraviolet A (UVA) light and treatment with riboflavin (vitamin B2) plus ultraviolet B (UVB) light, both targeting the nucleic acids of pathogens. The ability of the Mirasol PRT system to inactivate both pathogens and white blood cells has been previously described.¹ The technology uses a combination of riboflavin and UV light to induce irreversible lesions in the nucleic acids of pathogens and white blood cells (WBCs) to inhibit replication and function.²

Hence, PI techniques can be considered as a “Paradigm shift” in ensuring safer blood transfusion as PI uses a variety of physical, chemical, or photochemical methods to remove or inactivate that blood-borne pathogens, such as viruses, bacteria, and parasites in blood components or products. These PI methods include but not limited to solvent/detergent (S/D), nanofiltration and photochemical inactivation such as using Methylene Blue (MB), psoralens, or riboflavin.

Nowadays, research on PI technology for blood components (plasma and platelet) has made a great progress. Several inactivation methods including MB, Psoralens and Riboflavin can be chosen. These methods are targeting viral nucleic acids (NA) through photochemical inactivation. Methylene Blue (MB) is a phenothiazinium dye with a wide range of pathogens (bacteria, viruses and protozoa etc). Mirasol (Terumo BCT, USA), based on Riboflavin/UVB PI technology, has been used in a number of blood centers in many countries.³

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The advantages of PI includes (a) PI methods are particularly efficient to prevent transfusion-related bacterial infections (b) PI methods of PC globally reduce the risk of transfusion transmitted diseases and replace γ-irradiation for the prevention of GvHD. (c) Hemostatic efficacy of PI-treated PC appears to be maintained, although the CCI is lower when compared to untreated PC.

In a study conducted by Castrillo et al.,² the PC in PAS treated with the Mirasol system demonstrated minimal loss of platelet quality parameters. A minimal reduction in swirling was noted on the 7th day of storage, indicating that PC morphology was preserved in all units.

In vitro studies have shown that PI-treated PC has a raised activation level with increased metabolic parameters which needs to be further verified on a larger scale with extensive documentation. Activated fibrinogen receptor expression appears to be increased after PI, perhaps through a direct effect of PI on this integrin. These data relate mainly to the amotosalen/UVA technique and, to lesser extent, to the riboflavin/UV method.⁴

PI of blood components is a solid next step to eliminate the threat of transfusion transmissible disease in the blood supply. However, a well-balanced interplay of different aspects is needed to successfully
implement these worldwide. One of the most important criteria to ascertain include the cost- economics and the feasibility for wider application particularly in high demands and resource constrained setup as seen in developing countries. However, recent analyses have demonstrated that PRT-treated blood products may actually reduce the overall health care costs and the duration of hospital stay associated with post-transfusion care for some patients. Due to the ability of the Mirasol PRT treatment to inactivate all remaining leukocytes in a product that mediate GvHD, it can be used as an alternative to γ-irradiation thereby saving huge medical cost and ensuring increased patient comfort and satisfaction.6

In addition, the Mirasol PRT system, which is considered to be the gold standard to inactivate residual leukocytes in blood products, has been shown to prevent accumulation and secretion of most WBC–associated cytokines, with the potential to prevent leukocytes-mediated immunologic reactions in recipients.6

The Mirasol system is easy to use and does not require special training to operate the equipment. This technology can eliminate most of the residual risk of bacteria and also reduces the risk associated with a long list of transfusion-transmitted pathogens for which the blood supply is not screened.

More studies are required to fully understand the various mechanisms involved for PI and hence PI remains a challenge in the near future, which in turn, call’s for more concerted efforts for sustained and long term clinical benefits.

Acknowledgements

None.

Conflict of interest

The author declares no conflict of interest.

References


Citation: Das S. Pathogen inactivation: prospects and pitfalls. J Bacteriol Mycol Open Access. 2017;5(6):399–400. DOI: 10.15406/jbmoa.2017.05.00154