The monocyte activation test (MAT) is used to detect pyrogens in pharmaceutical products and serves as replacement of the rabbit pyrogen test. The peripheral blood mononuclear cell-based MAT assay requires the addition of serum to the medium and is performed with either fetal bovine serum (FBS) or human serum (HS). Since the capacity to detect non-endotoxin pyrogens (NEPs) in a sensitive manner is an important strength of MAT compared to the bacterial endotoxin test, the performance of the MAT using FBS and HS was compared using endotoxin and several NEPs. The MAT was more sensitive for endotoxin when FBS was used, however for most NEPs the MAT was more sensitive when performed in HS. Furthermore, heat-inactivation of FBS affected the performance of the MAT for endotoxin to some extent but not for the NEPs. Interestingly, heat-inactivation of HS led to an almost complete loss of reactivity towards endotoxin, reduced the response towards heat-killed Staphylococcus aureus and peptidoglycan, but had minor or no effects on the responses towards R848, flagellin, and Pam3CSK4. Product testing of a human blood-derived product in MAT using HS was beneficial since endotoxin spike recoveries were improved. This product is therefore currently batch released with the HS-based MAT assay. Overall, to guarantee optimal performance of MAT, heat-inactivated serum should be avoided. The HS-based MAT appears to be the first choice to replace the rabbit pyrogen test, while in some cases the FBS-based MAT may be favored.

1 Introduction

Pharmaceutical products intended for parenteral use must be shown to have pyrogenicity below a specified level to protect humans from “injection fever”, a marked rise in body temperature caused by injection/infusion of a drug product contaminated with pyrogens. Pyrogens are subdivided into endotoxins (lipopolysaccharides; LPS) derived from Gram-negative bacteria and non-endotoxin pyrogens (NEPs) such as peptidoglycans (PGN), flagellin and β-glucans derived from Gram-positive bacteria and fungi.

Historically, the rabbit pyrogen test (RPT) was used to determine pyrogenicity of injectable drugs for humans, including plasma-derived products and implantable medical devices. However, the RPT assay requires the use of animals, which should be avoided for ethical reasons where possible. Also, the responsiveness between humans and rabbits to pyrogens differs: Both species show a high reactivity to endotoxin, but humans usually show a higher reactivity to Gram-positive pyrogens than rabbits (Schindler et al., 2003).

The monocyte activation test (MAT) is an in vitro-based alternative to the RPT based on the capacity of human immune cells such as monocytes to respond to endotoxins and NEPs by production of pro-inflammatory cytokines like interleukin (IL)-6, IL-1β or tumor necrosis factor (TNF) (Dimarello, 2004; Kikkert et al., 2008). The current European Pharmacopeia (Ph. Eur.) chapter 2.6.30 (07/2017) states that “MAT is suitable, after product specific validation, as a replacement for the rabbit pyrogen test” (EDQM, 2020).

According to Ph. Eur. chapter 2.6.30, multiple cell sources can be used for the MAT assay, namely whole blood, peripher-
al blood mononuclear cells (PBMCs), or a monocyctic cell line as reviewed by Poole and colleagues (2003). The monocyctic cell line has the disadvantages that it represents just one genetic background, is derived from a tumor cell line, and functional stability over passaging has to be ensured. Whole blood or PBMCs may be either freshly isolated or thawed from cryopreserved material. Whole blood or PBMCs may be derived from a single donor or employed as pools from at least four donors (Solati et al., 2015; Koryakina et al., 2014). The requirement of four donors for pools is intended to include genetic variation in humans to the responsiveness to NEPs (Jaeger et al., 2015), while the responsiveness to endotoxins is conserved among humans (Copeland et al., 2005).

In this study, the MAT was performed with cryopreserved pooled PBMCs from four donors combined with an IL-6 ELISA as cytokine readout. Previous results from our and other laboratories have shown that a PBMC-based MAT assay has high sensitivity and high specificity (Hoffmann et al., 2005; Solati et al., 2015; Vipond et al., 2019; Etna et al., 2020).

According to the Ph. Eur. (EDQM, 2020), each new product has to go through a product-specific validation to determine that it does not interfere with the MAT test. This product-specific validation consists of 1) tests showing that the MAT detects endotoxins and NEPs when spiked in the product and 2) a test showing that the product does not interfere with the detection system (ELISA). Subsequently, release testing can be performed on the production process qualification (PPQ) batches (for registration) and on each production batch. During release testing according to method A (quantitative test) or method B (semi-quantitative test), as described in Ph. Eur. chapter 2.6.30 (EDQM, 2020), three dilutions of the product, which were determined during the product-specific validation, are tested with and without an endotoxin spike. The lowest dilution with a valid endotoxin spike recovery, between 50 and 200%, is used to calculate the amount of endotoxin equivalent units per milliliter (EEU/mL) for method A. If the responses to dilutions of a product are not comparable to the responses of standard endotoxin dilutions, then method B should be used. Method B is a pass/fail test, e.g., the contaminant concentration of the preparation should be below the contaminant limit concentration (CLC) to pass.

A PBMC-based MAT assay requires supplementation of the cell culture medium with serum. Serum contains proteins that aid the immune reaction triggered by pyrogens. The MAT can be performed with either fetal bovine serum (FBS) or human serum (HS) as serum source. Since the capacity to detect NEPs in a sensitive manner is an important asset of the MAT, we compared the performance of the MAT in FBS and HS to gain more insight into the detection of endotoxins and NEPs by the PBMC-based MAT using these different serum sources. This is especially relevant since results might support a MAT format without use of animal-derived materials. In addition, we studied the effect of heat-inactivation of the serum on the response to endotoxin and NEPs and the effect of serum source on the recovery of the endotoxin spikes in a specific human blood-derived product.

2 Materials and methods

Pyrogens
Endotoxin Standard Biological Reference Preparation (BRP) batch 5 (EDQM, Council of Europe, Strasbourg, France) was used to prepare the endotoxin standard curve. A stock solution of 2000 EU/mL was prepared in endotoxin-free water, aliquoted (50 µL), and stored at -80°C. The endotoxin stock solution was diluted further in complete medium to the required endotoxin concentrations. Heat-killed S. aureus (HKSA), flagellin (S. typhimurium), Pam3CSK4, R848 and PGN (S. aureus) were all obtained from Invivogen (San Diego, CA, USA) and aliquoted and stored according to the manufacturer’s instructions. The absence of endotoxins in these NEP preparations was claimed by the manufacturer and reconfirmed in-house for HKSA, flagellin and PGN using the Limulus Amebocyte Lysate (LAL) test (Pam3CSK4 and R848 are not suitable for testing in the LAL).

Of note, the endotoxin and NEP concentrations stated throughout this study are final concentrations in the well (after addition of cells to the well). Concentration in the sample can be calculated by multiplying the stated concentrations by 2 (since samples and cells are mixed 1:1).

MAT execution: Cell culture
The MAT was performed using the MAT Cell Set (Sanquin Reagents, Amsterdam, The Netherlands) following the manufacturer’s instructions with adaptations as described: In short, a vial containing 5x10^6 (± 1x10^6) cryopreserved PBMCs (1 mL) was thawed in a water bath at 37°C until only a little clump of ice remained. The cell suspension was subsequently transferred to a new tube and diluted with 10 mL Iscove’s Modified Dulbecco’s Medium (IMDM; Lonza, Verviers, Belgium) supplemented with either 5% FBS or HS (both from Biowest, Nuaillé, France). Where indicated, FBS or HS was heat-inactivated at 56°C for 45 min before use. In the MAT cell culture plate, 100 µL cell suspension was added to 100 µL sample (tested in triplicate or quadruplicate where indicated). In case of spiking experiments, 5 µL supplemented IMDM was added to the thawed cell suspension, and 50 µL cell suspension was added to 100 µL sample plus 50 µL endotoxin spike solution or 50 µL supplemented IMDM (for unspiked samples). In both cases, each well contained approximately 45,000 cells. Subsequently, the 96-well cell culture plate was incubated in a humidified incubator at 37°C in the presence of 5% CO₂ for 18-24 h.

MAT execution: IL-6 ELISA
Cell culture supernatants were harvested and analyzed in 1:5 dilution, except if indicated otherwise, for the presence of IL-6 using a commercially available enzyme-linked immunosorbent assay (ELISA) kit (PeliKine Compact™ human ELISA kit, Sanquin, Amsterdam, The Netherlands) following the manufacturer’s instructions. The ODs at 450 nm after subtraction of the background OD at 540 nm are depicted.

Drug product testing with the MAT
To exclude interference between the test and the product being
Fig. 1: Comparison of MAT performed in FBS and HS
PBMCs were stimulated overnight with endotoxin (LPS), HKSA, Pam3CSK4, PGN, flagellin and R848 in a 2-fold concentration range using either FBS or HS as serum source. An IL-6 ELISA was performed on the culture supernatants; supernatants of R848 were diluted 1:25 for IL-6 determination; for all other conditions, supernatants were tested in 1:5 dilution. (A) Representative graphs are depicted; results are shown as mean OD ± SD of triplicate measurements. (B) Area under the curve (AUC) of fitted pyrogen curves of all experiments is calculated and depicted as mean ± SD. Endotoxin (LPS) (n = 4), HKSA (n = 4), Pam3CSK4 (n = 4), PGN (n = 3), flagellin (n = 5) and R848 (n = 4). Paired t-test was performed. *p < 0.05, ***p < 0.001.
tested, the MAT has to be validated for that product according to the pharmacopeia guidelines. For this, endotoxin is spiked into uncontaminated batches of the product and tested in the MAT. The product is considered free from interfering factors if the mean recovery of the added endotoxin is in the range of 50-200%. During these experiments, all samples were tested in quadruplicate and 50 µL cell suspension was added to 100 µL sample and 50 µL endotoxin spike or 50 µL complete medium (for unspiked samples). The endotoxin spike was 0.02 EU/mL for the FBS-based MAT or 0.16 EU/mL for the HS-based MAT. In addition, an endotoxin standard curve was included on the same plate in quadruplicate, ranging from 0.005 to 0.08 EU/mL for FBS-based MAT and 0.02 to 0.32 EU/mL for the HS-based MAT.

Statistical analysis
GraphPad Prism 8.0.2. was used for the analysis. Area under the curve (AUC) of nonlinear fitted regression lines ([agonist] vs. response-variable slope) were calculated. Paired t-test was used to compare the AUC of the two serum sources. Unpaired t-test with correction for multiple comparison with Holm-Sidak method was performed to analyze the effect of HS in FBS-based MAT on detection of endotoxin.

3 Results

3.1 Comparison of MAT responses to endotoxin and NEPs using FBS and HS as serum source
To investigate if the reactivity of the MAT is different for endotoxin and a selection of NEPs using FBS or HS as serum source, we performed the MAT with medium containing either 5% FBS or 5% HS. Sensitivity towards endotoxin was lower in the HS-based MAT compared to the FBS-based MAT. The limit of detection (LOD) for endotoxin using FBS was ≤ 0.01 and for HS ≤ 0.03 EU/mL per well (Fig. 1A). The HS-based MAT was more sensitive towards HKSA (a ligand for toll-like receptor-2 (TLR-2)), Pam3CSK4 (a synthetic mimic of bacterial lipopeptides (TLR-2/1 ligand)), and peptidoglycan (PGN; a NOD1/2 ligand from S. aureus) in comparison to the FBS-based MAT. In case of flagellin (TLR5 ligand from gram-negative Salmonella typhimurium) and imidazoquinoline compound R848 (TLR7/8 ligand), we did not observe a significant difference in reactivity between MAT using FBS or HS. The observed differences between performing MAT using FBS or HS were not dependent on the PBMC pool, batch of HS or batch of FBS (data not shown). Overall, we observed that the MAT is more sensitive for endotoxin when performed with FBS instead of HS, while for the NEPs the sensitivity was either lower or the same when performed with FBS compared to HS.

3.2 Effect of heat-inactivation of serum on detection of endotoxin and NEPs by MAT
Heat-inactivation of serum is usually performed to inactivate complement (Solits et al., 1979). However, heating of serum also reduces or modifies serum growth factors and is often performed without evidence of a beneficial effect (Rahman et al., 2011). To study if the reactivity of the MAT is affected by heat-inactivation of the serum, we compared the responses in MAT using untreated or heat-activated FBS or HS.

Heat-inactivation of FBS reduced the reactivity of the MAT towards endotoxin (Fig. 2A,B) but not towards NEPs. Heat-inactivation of HS had a more pronounced effect compared to heat-inactivation of FBS (Fig. 2C,D). It resulted in an almost complete loss of reactivity of MAT towards endotoxin and had varying effects on reactivity towards NEPs. The reactivity of the MAT to Pam3CSK4 was not affected by heat-inactivation of HS, the reactivity to R848 and flagellin was significantly higher, but the response towards HKSA and PGN was significantly lower using heat-inactivated HS. Overall, heat-inactivation of HS had differential effects on the responses of the MAT to pyrogens.

3.3 Interference of HS in an FBS-based MAT
Since the reactivity towards endotoxin was lower in the HS-based MAT compared to the FBS-based MAT, we wondered if HS contains either lower amounts of stimulatory constituents or higher amounts of inhibitory constituents affecting the endotoxin response. Therefore, we performed an FBS-based MAT in IMDM with 5% FBS using a fixed amount of endotoxin (0.08 EU/mL) and added increasing amounts of HS to investigate if this affected the response towards endotoxin. To correct for the higher total concentration of serum, we added the same amount of additional FBS to controls containing the same amount of endotoxin. The experiments were performed with serum that had not been heat-inactivated.

The addition of HS caused a significant inhibition of the response towards endotoxin, while this was not the case for addition of FBS (Fig. 3). The inhibition was already observed at 0.16 % HS, but was more pronounced at higher HS concentrations. This indicates that HS contains constituents that can inhibit the response to endotoxin in an FBS-based MAT.

3.4 Effect of serum source during drug product testing
In previous studies, it was observed that certain plasma-derived products yielded invalid recoveries of the endotoxin spike that was added to the products, making it complex to test these products in the MAT (method A or B of Ph. Eur. 2.6.30) (Perdomo-Morales et al., 2011). Here a blood-derived product was tested in the MAT, using either FBS or HS as serum source, to assess the effects on endotoxin recovery. The maximum valid dilution of a product is determined by dividing the contaminant limit by the LOD.

With the FBS-based MAT, the LOD was 0.008 EU/mL, and this resulted in a maximum valid dilution (MVD) for the product of 625-fold. The product was tested at 150-, 300- and 600-fold dilutions; however, the 150-fold dilution did not yield a valid endotoxin recovery (< 50%) (Fig. 4). The 300-fold product dilution yielded recoveries around 50% and the 600-fold around 60%. This resulted in many repeats of the routine MAT assay due to invalid endotoxin spike recoveries.

The same blood-derived product was also tested with the HS-based MAT. Since the LOD of this MAT was determined to
Fig. 2: Effect of heat-inactivation of serum on MAT
PBMCs were stimulated overnight with endotoxin (LPS), HKSA, Pam3CSK4, PGN, flagellin and R848 in 2-fold concentration range. The effect of heat-inactivation of FBS (A,B) and HS (C,D) on the MAT response was tested. An IL-6 ELISA was performed on the culture supernatants; supernatants of R848 were diluted 1:25 for IL-6 determination; for all other conditions, supernatants were tested in 1:5 dilution. (A,C) Representative graphs are depicted, results are shown as mean OD ± SD of triplicate measurements. (B,D) Area under the curve (AUC) of fitted pyrogen curves of all experiments is calculated and depicted as mean ± SD. Endotoxin (LPS) (n = 4), HKSA (n = 4), Pam3CSK4 (n = 4), PGN (n = 3), flagellin (n = 5) and R848 (n = 4). Paired t-test was performed. *p < 0.05, **p < 0.01, ***p < 0.001.
Fig. 2: Effect of heat-inactivation of serum on MAT (cont.)

PBMCs were stimulated overnight with endotoxin (LPS), HKSA, Pam3CSK4, PGN, flagellin and R848 in 2-fold concentration range. The effect of heat-inactivation of FBS (A,B) and HS (C,D) on the MAT response was tested. An IL-6 ELISA was performed on the culture supernatants; supernatants of R848 were diluted 1:25 for IL-6 determination; for all other conditions, supernatants were tested in 1:5 dilution. (A,C) Representative graphs are depicted, results are shown as mean OD ± SD of triplicate measurements. (B,D) Area under the curve (AUC) of fitted pyrogen curves of all experiments is calculated and depicted as mean ± SD. Endotoxin (LPS) (n = 4), HKSA (n = 4), Pam3CSK4 (n = 4), PGN (n = 3), flagellin (n = 5) and R848 (n = 4). Paired t-test was performed. *p < 0.05, ** p < 0.01, ***p < 0.001.
be 0.03 EU/mL, the maximum valid dilution of the product was only 166-fold. Therefore, the product was tested at lower dilutions, namely 31.2-, 62.5- and 125-fold dilutions. All 3 product dilutions consistently yielded valid endotoxin recovery values between 50-200% (Fig. 4A). After product-specific validation, the routine measurements were performed with HS-based MAT, and the data from 20 batches showed the same result: although the MAT assay based on HS serum requires higher product concentrations, the endotoxin recoveries of an endotoxin spike were consistently good (Fig. 4B).

### 4 Discussion

In this study, we compared the performance of the MAT using either FBS or HS as serum source. Except for a slightly lower sensitivity for endotoxin, the MAT was more reactive towards most NEPs when performed with HS. Heat-inactivation of FBS affected the performance of the MAT for endotoxin to some extent but not for the NEPs. However, heat-inactivation of HS led to an almost complete loss of reactivity towards endotoxin and reduced reactivity in the response towards HKSA and PGN, while having minor or no effects on the responses towards R848, flagellin and Pam3CSK4. Moreover, batch release testing of a blood-derived product in MAT using HS was beneficial since endotoxin spike recoveries were valid at lower product dilutions.

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**Fig. 3: Interference of HS on endotoxin-induced response in an FBS-based MAT**

PBMCs were stimulated overnight with 0.08 EU/mL endotoxin in medium containing 5% FBS. HS or FBS was added in graded doses as indicated. An IL-6 ELISA was performed on the culture supernatants; OD values were normalized to samples with 5% FBS. Data of 5 independent experiments are combined and represented as mean ± SD. Unpaired t-test with correction for multiple comparison with Holm-Sidak method was performed. **p < 0.01, ***p < 0.001.

**Fig. 4: Difference in product interference on endotoxin recovery in FBS- and HS-based MAT**

(A) Endotoxin recovery of three batches of a blood-derived product with FBS- and HS-based MAT. Mean of 4 replicates is depicted. (B) Endotoxin spike recovery of 20 batches tested with FBS-based MAT and 20 different batches tested with HS-based MAT. The mean of each sample quadruplicate is shown as an individual point. The overall error bar is the mean of the samples ± SD. Dotted lines depict the border for valid recoveries of the spike (> 50 and < 200).
Several co-factors are required for the stimulation of TLR4 by endotoxin including LPS-binding protein (LBP), CD14 and MD-2 (as reviewed in Raetz and Whitfield, 2002; Park and Lee, 2013). MD-2 forms a receptor complex with TLR4 at the membrane and is required for the recognition of LPS molecules by the TLR4/MD-2 complex and subsequent signaling events. LBP is a soluble protein present in serum and is necessary for the extraction of LPS from the bacterial surface or from LPS aggregates. LBP transfers the LPS molecule to CD14, which presents the LPS molecule to the TLR4/MD-2 complex. CD14 is present as a GPI-anchored membrane protein on monocytes and also in soluble form in serum.

Endotoxin was the only pyrogen tested that gave a higher reactivity in FBS compared to HS. Differences in LBP concentration between FBS and HS might explain the difference in reactivity. However, since additional HS in an FBS-based MAT inhibited the response to endotoxin, this indicates that (in addition) inhibitory components are present in HS which might (partly) lower the response to endotoxin in HS-based MAT. It has been described that serum lipoproteins in HS, e.g., LDL and VLDL, which can inhibit the response to endotoxin (as reviewed by Berbee et al., 2005; Wendel et al., 2007), are absent or present in lower amounts in FBS (Haylett and Moore, 2002; Forte et al., 1981). Furthermore, the presence of anti-LPS antibodies in human serum, possibly blocking the binding to the TLR4/MD-2 complex, can also not be ruled out (Fomsgaard et al., 1987; Nys et al., 1996).

Heat-inactivation of FBS led to a reduced response towards endotoxin, while heat-inactivation of HS almost completely blocked the response towards endotoxin, demonstrating that heat-inactivation of serum is neither beneficial nor advisable for detection of endotoxin in the MAT. The effect of the heat-inactivation might be explained by the heat-sensitivity of LBP (Meszaros et al., 1995).

The reactivity to the different NEPs in the MAT was much higher or comparable when performed with HS compared to FBS. Especially the response towards HKSA and PGN increased substantially. The response towards Pam3CSK4, a synthetic lipopeptide that mimics the acylated amino terminus of bacterial lipoproteins (a constituent of the cell wall of Gram-positive and Gram-negative bacteria), was also enhanced in the HS-based MAT.

Heat-inactivation of HS had differential effects for the different NEPs tested. The most profound effect was the lowered response towards HKSA and PGN. Altogether, heat-inactivation of HS on responses towards endotoxin, HKSA and PGN impacted the overall performance of the MAT in HS far more than the slight positive effect of heat-inactivation of HS observed for R848 and flagellin.

Drug product testing in the MAT requires that endotoxin spike recoveries in the product are between 50% and 200%. Some plasma-derived products show high interference in the MAT, rendering the assay unsuitable as a pyrogen test due to invalid spike recovery (Perdomo-Morales et al., 2011). In some cases, especially with a highly sensitive MAT (low LOD), interference can be overcome by dilution of the product close to the MVD. Although our HS-based MAT had a lower MVD due to a higher LOD for endotoxin, testing of a blood-derived product in this assay resulted in valid spike recovery at 10-20-fold lower product dilutions compared to the FBS-based MAT (31.2-fold vs 300-600-fold diluted). A possible explanation for obtaining valid spike recoveries at lower product dilutions using the HS-based MAT compared to the FBS-based MAT could be that an inhibitory component of HS provides a form of basal inhibition towards endotoxin in the assay (as indicated by a higher LOD). This may limit further inhibition of any possible inhibiting component in the drug product, resulting in improved spike recovery. This is supported by our finding that adding HS to the FBS-based MAT results in interference. The HS-based MAT has been routinely used for releasing batches of this specific blood-derived medicinal product since 2018.

In conclusion, the MAT is more sensitive for endotoxin detection when FBS is used as serum source; however, the MAT is more sensitive for most NEPs when performed in HS. Further, we show that the HS-based MAT is preferred for certain blood-derived products. The HS-based MAT is therefore considered to be the first choice to replace the rabbit pyrogen test, since the detection of certain NEPs is superior compared to FBS. HS also better reflects the human immune system and, importantly, avoids the use of animal-derived FBS, which is under debate as a cell culture supplement owing to animal welfare concerns and compliance with the 3R principle (van der Valk et al., 2018).

Recently, several papers have described the use of MAT for pyrogen testing in vaccines (Vipond et al., 2019; Etna et al., 2020; Rossi et al., 2020). Vaccines are complex products since they differ in formulations (e.g., type of antigens, adjuvant, excipient and stabilizer), and some vaccines have inherent pyrogenic activities while others do not. For vaccines with pyrogenic activity, the amount of pyrogenicity should be balanced: too much immune activation will lead to more severe side-effects (e.g., fever), too little will not provide protection. Especially for vaccines with pyrogenic activity, the MAT can serve as a release test to determine the consistency between batches. The FBS-based MAT however could be more suitable for testing certain vaccines, especially those against diseases for which the HS donors may already have antibodies, which could interfere in the MAT assay, e.g., by binding to the vaccine. Finally, to guarantee optimal performance of the MAT, heat-inactivation of serum should be avoided.

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Conflict of interest
Sanquin Diagnostics performs MAT testing for customers. Sanquin Reagents produces and distributes MAT cell kits.

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