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COMPARISON OF CELLULAR ASSAYS FOR TOLL-LIKE RECEPTOR ACTIVATION AND DEVELOPMENT OF A SPECIES-SPECIFIC REPORTER CELL LINE FOR CATTLE

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Abstract
Pattern recognition receptors (PRRs) form the first defence line of the immune system. Toll-like receptors (TLRs) are the best known and the most examined of the PR receptors. In the last several years, TLRs have been studied in different ways resulting in a lot of new insights into the function and signalling pathways of these receptors. However, as significant differences exist between species, it is also necessary to develop and adapt these tools for economically important farm animals, especially if one wants to investigate individual combinations of the TLRs and their specific ligands, because of the complex network in immune signalling resulting in interference or synergy with one another. It is increasingly important to discover new vaccine adjuvants engaging the innate immune system for production animals to reduce the impact of disease and antibiotic use. This work shows a species-specific cell-based assay adapted for the analysis of single PRRs. For this purpose, HEK (human embryonic kidney) 293T cells were stably transfected with the NF-κB-inducible reporter gene secreted embryonic alkaline phosphatase (SEAP) together with bovine TLR2. We compared the SEAP response with an existing firefly luciferase NF-κB reporter assay for correlation with IL-8 production in the context of bovine TLR2. A dose dependent NF-κB response was detected upon FSL-1 stimulation using both methods with good correlation to IL-8 protein secretion (Spearman’s r=0.9879; p<0.0001 in case of SEAP and r=0.7939; p=0.0088 with the luciferase assay). TLR2 activation was detected at lower FSL-1 concentrations by the SEAP assay compared to IL-8 secretion at the same time point. The luciferase assay produced very high non-specific background for all ligand concentrations. Of all three assays tested we found the bovine specific TLR2 NF-κB SEAP reporter assay to be the most convenient with fewer steps and delivered results in the shortest time. The developed reporter cell line would lend well to rapid, high throughput TLR ligand screening for cattle.

Keywords
Bovine TLR2, reporter assay, SEAP, luciferase assay, CXCL-8, IL-8

Highlights
• To study individual TLR activation, it is most convenient to use cell lines expressing only the receptor in question
• We developed a bovine TLR2-expressing SEAP reporter cell line
• We compared luciferase and SEAP assays, with IL-8 detection as control
• Both methods detect TLR-stimulation, however, the luciferase assay gives high non-specific background signal in the presence of the reporter gene
• Of all tested methods, stable SEAP reporter cell lines were the most convenient to use
1. Introduction

The innate immune system is the first defence line against pathogenic microbes. It can be classified into two parts: the afferent (sensing) and the efferent (effector) system. Pattern recognition receptors (PRRs) like toll-like receptors (TLRs), dectins, CD14 or NOD-like receptors (NLRs) belong to the afferent system together with the humoral components, e.g. lipopolysaccharide binding protein (LBP), collectins and C3b (Beutler, 2004). PRRs are involved in early detection of specific pathogens because they recognize highly conserved and class-specific motifs called microbe associated molecular patterns (MAMPs). The best characterized class of the PRRs is the evolutionary conserved TLR family, first recognized in Drosophila (Lemaître et al., 1996). In most mammalian species 10 TLRs have been described to date and they differ from each other in expression patterns, ligand specificity and in the target genes they induce (Jungi et al., 2011). TLRs control the activation of innate immunity and induce activities against invading pathogens by the production of inflammatory cytokines. They link innate and adaptive immunity through the induction of antigen-presentation and co-stimulatory molecules.

TLRs are expressed on a variety of immune and non-immune related cells (Akira et al., 2006). Targeting TLRs with specific ligands shows great potential for application as vaccine adjuvants (Bode et al., 2011; Mizel and Bates, 2010). Whereas ligand specificities for most TLRs are fairly described, novel interactions are continuously being discovered (Salazar Gonzalez et al., 2014), and no known ligand has been described for TLR10 in any species. Despite the fact that TLRs are highly conserved between mammals, most available information on TLR function was generated in murine or human systems and cannot be readily extrapolated to other, economically important farm or companion animal species (Metcalf et al., 2014). The identification of new vaccine targets/adjuvant, immunomodulatory targets is becoming an increasing aim of research, given the increasing abuse and subsequent ban of antimicrobial compounds for use in farm animals (Toutain et al., 2016). For these reasons, several methods have been developed to screen potential TLR-ligands in a fast and reproducible manner. The majority of these assays rely on the ability of a known ligand stimulating transcription factors such as the nuclear factor κ-light-chain-enhancer of activated B cells (NF-κB), activating protein-1 (AP-1) or interferon regulatory factor (IRF) family (Kawai and Akira, 2006) in a TLR dependent manner. To do so, cellular assays involving either primary immune cell subsets or cell lines expressing specific TLRs are normally employed. Testing TLR activation in vitro with established cell lines is easier to standardise and effectively reduces the use of experimental animals. Although such lines may not be as biologically relevant as primary cells, ligand specificity of the receptor in question can be tested and compared regardless of which downstream events are activated. The decision of cell lines to be used has to take endogenous TLR expression, need for potential co-receptors and unresponsiveness to standard ligands, such as LPS into account. Transfection of such cell lines with specific TLRs allows for end products of activation (transcription factors, cytokines) to be detected and quantified.

Indeed, several colorimetric reporter gene assays have been developed allowing for the detection of transcription factor activation. Luciferase-based assays, such as dual luciferase assays are among the most popular reporter assays used for pattern recognition receptor activation (Ling et al., 2012). Here, TLR-specific activation of NF-κB is measured using a reporter gene coding for the firefly (Photinus pyralis) luciferase, which is measured against the activity of the sea pansy (Renilla reniformis) luciferase using a luminometer. Another popular assay utilizes secreted embryonic alkaline phosphatase (SEAP) as a reporter gene. The application of SEAP as a reporter for TLR activation was introduced by Burger-Kentischer et al. (2010) with the aim of substituting the limulus amoebocyte lysate (LAL) test for endotoxin and pyrogen detection. The advantage of this assay is that it does not rely on a luminometer to detect activation of NF-κB, but rather on a colorimetric change based on the activity of alkaline phosphatase, which can be either detected by eye or quantified using a conventional ELISA plate reader. The SEAP assay has been further developed and stable reporter cell lines expressing human or murine TLRs are commercially available, along with substrate containing cell culture media enabling
real-time quantification of SEAP activity; however, the assay has to be carefully controlled to avoid detection of endogenous alkaline phosphatase activity from cells or culture media.

Lastly, activation of transcription factors can also be assessed by analysing production of downstream molecules, such as secreted cytokines/chemokines. This method has the advantage that it can be easier correlated with the response in primary cells, but requires a direct correlation between transcription factor activation and cytokine/chemokine protein production.

In the present study, we compared the three different test systems mentioned above using luciferase, SEAP and IL-8 production for bovine TLR2, for which we already had identified species-specific ligand differences (Willcocks et al., 2013).

2. Materials and methods

2.1 Generation of SEAP reporter cell line

All cell lines used in these experiments were generated using the human embryonic kidney (HEK) 293T cells kindly provided by Dr. Bradley Cobb (The Royal Veterinary College, University of London) and were negative for endogenous TLR1, 2 and 6 expression as established by RT-PCR and flow cytometry (data not shown). The cells were maintained in complete 293T medium; DMEM supplemented with GlutaMAX (Gibco, UK), 10% foetal bovine serum (FBS; Sigma), 100 IU/ml–100 μg/ml penicillin–streptomycin (Gibco) and 1mM sodium-pyruvate (Sigma). The vector pNifty2-SEAP (Invivogen) was transfected into 293T cells using TurboFect reagent (Thermo Scientific) according to manufacturer’s protocol for six-well plates. Transfectants (293T/SEAP) were clonally selected according to the method described by Burger-Kentischer et al. (2010) using 100 μg/ml Zeocin (Invivogen) and maintained in complete selective medium afterwards. Endogenous SEAP activity was found to be minimal in response to a panel of TLR ligands for 293T and 293T/SEAP cells (data not shown).

2.2 Generation of bovine TLR2-expressing 293T and 293T/SEAP reporter cell lines

The bovine TLR2 gene (Willcocks et al., 2013) was cloned into the mammalian expression vector pTracer-CMV/Bsd (Thermo Scientific) and transfected into 293T and 293T/SEAP reporter cells, followed by clonal selection using 10 μg/ml Blasticidin (Sigma-Aldrich). TLR2 expression was confirmed by flow cytometry (data not shown). Both cell lines were maintained in complete selective medium (containing Zeocin and Blasticidin for 293T/SEAP/TLR2 and Blasticidin only for 293T/TLR2 cell lines).

2.3 Stimulation assay of 293T/SEAP/TLR2 reporter cells

293T/SEAP/TLR2 cells were seeded into a 96-well plate at 2x10⁴, at 3x10⁴ and at 4x10⁴ cell/well densities in 200 µl/well complete selective media (using 12 wells of each concentration, Fig. 1A), and incubated in a 37 °C 5% CO₂ humidified incubator. After 24 hours, culture media was exchanged for stimulation media (DMEM-Glutamax, supplemented with 2% FBS and 1 mM sodium-pyruvate) and 1, 10, 100, 500 or 1000 ng/ml FSL-1 (Invivogen), 200 ng/ml phorbol 12-myristate 13-acetate (PMA, Sigma) or no stimulant in duplicate wells. FSL-1 is a synthetic lipoprotein derived from Mycoplasma salivarium, used as a specific TLR2 agonist and PMA is a non-specific NF-κB stimulator, added as positive control with optimal stimulation levels previously determined (data not shown). Following an additional 24 h incubation, supernatants were collected and frozen at -20 °C until measurements of SEAP activity and IL-8 production. SEAP activity was measured in a 96-well clear flat-bottom cell culture plate; 10 µl of each sample supernatant was added to 200 µl prewarmed QuantiBlue reagent (Invivogen) and incubated at 37 °C. After 24 hours, optical densities at 635 nm were measured using a Tecan Infinite M200 Pro plate reader. IL-8 production was measured with Quantikine Human
CXCL8/IL-8 ELISA kit (R&D systems) according to the protocol suggested by the manufacturer. Fifty µl of each cell supernatant was used as sample and absorbance was measured at 450 nm.

2.4 Stimulation assay of 293T/TLR2 cells
293T and 293T/TLR2 cells were seeded at 10^5 cell/ml density in 4 ml complete media/well, into 3 wells each on a 6-well plate. After 24 hours incubation, one well of each cell line was transfected with 500 ng luciferase reporter plasmid NF-κB -Luc (pGluc, Patterson et al., 2014), 4 µg pNifty2-SEAP or mock (4 µg irrelevant plasmid vector). DNA amounts were previously optimised. After an additional 24 hours, the supernatants were discarded, the cells lifted and reseeded into 12 wells on a 96-well plate at 4x10^4/well density (Fig. 1B). Stimulation media was prepared with 125, 250, 500 or 1000 ng/ml FSL-1, 200 ng/ml PMA or no stimulant. After 24 hours of stimulation, luciferase activity from mock transfected cells and cells transfected with luciferase reporter gene was measured as described by Patterson et al. (2014) and supernatants were frozen until further processing. Human IL-8 levels and SEAP activity were assayed as described above.

Figure 1. Assay layouts for stimulation experiments using reporter cell line 293T/SEAP/TLR2 (A) and transient transfections (B). In plate A, cells were seeded at 2x10^4/well (light orange), 3x10^4/well (orange) and 4x10^4/well (dark orange) densities. PMA was used as positive control at 200 ng/ml in wells outlined purple. In replicates highlighted using blue, TLR-2 ligand FSL-1 was administered in increasing concentrations. Cells in row “A” were left unstimulated. SEAP activity and IL-8 production was measured from every well. 293T (orange) and 293T/TLR2 (red) cells, transfected with reporter plasmid pGluc (rows labelled “G”), pNifty-2-SEAP (rows labelled “N”) or mock (labelled “M”) were seeded into plate B at a density of 4x10^4/well. Cells in columns 1 and 2 were untreated. Positive control (purple wells) were stimulated with 200 ng/ml PMA, blue-rimmed wells with increasing amounts of FSL-1.

2.5 Statistical analyses
Calculations were performed using Prism 6 for Windows (GraphPad Software Inc). Optical density (OD) values imported from ELISA results were extrapolated using four-parameter logistic regression.
Invalid results were excluded from further analyses. OD values from SEAP assay were used as results in further comparisons. Signals measured throughout 10 seconds of the luciferase assay plate were averaged separately for each well. Multiple unpaired t-tests with Bonferroni correction were used to compare responses within measurement groups between stimulant doses. Spearman’s correlation was computed and Deming’s regression was plotted to compare methods separately for biological repeats.

3. Results and discussion
3.1. Responses of stable reporter cells
In order to develop a fast screening assay to identify farm animal specific TLR ligands, we created a stable boTLR2-expressing 293T SEAP reporter cell line, by inserting the reporter gene first, followed by the introduction of bovine TLR2. The 293T/SEAP/TLR2 cells responded to the synthetic lipopeptide FSL-1 in a dose-dependent manner (Fig. 2A and 2B). Within the combined data from three repeats, the lowest detectable, although not significant, specific ligand dose was 1 ng/ml for all seeding densities. Cells seeded at 4x10⁴/well gave the most pronounced OD differences for each FSL-1 concentration analysed. In contrast, when cells were seeded at 2x10⁴ or 3x10⁴/well, we observed decreasing signal detection above 500 ng/ml FSL-1 dose, probably because lower amounts of cells reach maximum binding capacity at lower ligand concentrations, resulting in a lower overall SEAP activity. Using the same supernatant, we detected the first significant increase in IL-8 concentration when stimulated with 500 ng/ml FSL-1.

Owing to the fact that two different substances (SEAP vs IL-8) were analysed, we could not apply Bland-Altman plot and Passing-Bablok regression, which are normally used to compare different measurement methods. Instead, a Spearman’s correlation between SEAP and IL-8 was used and shown to be significant (r=0.9596, p<0.0001). To demonstrate this correlation, a dot plot fitted with a Deming line was drawn (Fig. 2C). The regression line shows a considerable shift along the X axis (depicting SEAP values) towards the right (X=0.6427 when Y=0). Given that IL-8 values started to rise only at higher FSL-1 concentration, this shift was expected; however, at higher concentrations, the correlation remains strong between SEAP and IL-8 production. Optical densities measured in supernatants from cells stimulated non-specifically with PMA were at least 2.7-times higher (SEAP assay) or 17-times higher (ELISA) than those measured in non-stimulated cell media, validating the assays.

3.2. Stimulations of transient reporter cells
As the luciferase based reporter assay is based on a transient transfection system, we next compared the SEAP reporter system in a transient form with luciferase system and IL-8 ELISA. To do so, we generated bovine TLR2-expressing 293T cell clone (293T/TLR2), which were subsequently transiently transfected with luciferase or SEAP reporter genes, or mock transfected. IL-8 production was measured in all 293T/TLR2 cell clones. The results are presented in Fig. 3. Similar to the data obtained using a stable transfection system, we measured a dose dependent ligand response in the cells expressing TLR2 and transfected with the SEAP reporter gene, but not in cells lacking either the TLR, the reporter gene or both (Fig. 3A). In contrast, transient transfection with the luciferase reporter gene resulted in a relatively high background in cells containing the reporter gene but lacking TLR2 expression (Fig. 3B). This activity did not seem to be influenced by exposure to any of the FSL-1 concentrations used (Spearman’s r=0.7; p=0.2333). Lack of measurable IL-8 production in these cells (apart from the positive control) confirmed that this background reporter activity is unlikely to be NF-κB activation resulting from either endogenous TLR expression in undetectable levels or by non-specific means. As such, IL-8 levels were incomparable to those measured in cells where luciferase and also TLR2 were present (Fig. 3C). ELISA results were valid in all 293T/TLR2 replicates and IL-8 production was induced in all TLR2 expressing cell clones (Fig. 3D). Interestingly, the highest amount of IL-8
production was seen in mock transfected 293T/TLR2 cells. One potential explanation for this observation is competition for NF-κB between binding sites. The high amount of vector DNA containing NF-κB responsive elements introduced into transfected cells (500 ng/well in case of pGluc and 4 µg/well in case of pNifty-2) can engage a substantial amount of the transcription factor before translocation to the nucleus, while in mock transfected cells, all of these molecules can be employed in pro-inflammatory cytokine stimulation. In 293T/TLR2 cells, both reporter assays correlated well with IL-8 production (Spearman’s $r = 0.9879$ and $P < 0.0001$ in case of SEAP; and $r = 0.7939$ and $P = 0.0088$ with the luciferase assay within one biological repeat).

**Figure 2.** Results of reporter cell line stimulation and comparison of SEAP and IL-8 assays. Means of replicate values are indicated with SEM. (A) Dose-dependent SEAP-activity in 293T/SEAP/TLR2 cells at different seeding densities, the first significant increment indicated in each group. (B) Dose-dependent IL-8 production in 293T/SEAP/TLR2 cells at different seeding densities, the first significant increment indicated in each group. (C) Correlation of optical densities measured in SEAP and ELISA assays of the same samples fitted with Deming’s regression line and equation. Levels of significance: *$p<0.05$; **$p<0.01$; ***$p<0.001$.
3.3. Conclusions

Our results show that the SEAP based reporter system proved to be the most reliable and sensitive assay to detect bovine TLR dependent stimulation of NF-κB, and showed a good correlation with the resulting IL-8 production. Of all three assays tested we found the bovine specific TLR2 NF-κB SEAP reporter assay the most convenient with fewer steps and shortest duration. The developed reporter cell line would lend well to rapid, high-throughput bovine TLR2 ligand screening intended for supplementing vaccine adjuvant design strategies. Our findings extend recently published work by Tahoun et al. (2015). Similarly to previous findings (Willcocks et al., 2013), the TIR domain of the bovine receptor initiated adequate response in human cells in all three tested assays. In the present system, this response was already visible at 1 ng/ml, reaching significant differences at 10 ng/ml of specific stimulant doses in the SEAP assay when a stable reporter cell line was used. It was also observed that in stable reporter cell
lines, a higher maximum signal was detected than in TLR2-expressing cells transiently transfected with SEAP gene (averaging at an OD of 1.27 for stable and at 0.76 for transient cells), indicating the benefits of creating stable reporter cell lines for farm animal TLRs, as it already exists for human and murine systems. This way, there is no need for further transfections and the assay can be performed in a few simple steps, making it the assay of choice for small-scale laboratory experiments and also for high throughput automated screening. In contrast, and as experienced before, a high background was detected in the luciferase assay, based on non-specific luciferase activity in 293T cells lacking TLR2, despite the fact that stimulation resulted in a marked increase of the signal. This phenomenon can be more problematic when detecting lower doses of the agonist or when applied for ligand screening, where effective stimulant concentrations are unknown. In most cases, assessment of IL-8 production provided a useful control for TLR-ligand screening methods; however, cytokine production was undetectable particularly at lower ligand concentrations. It also has to be kept in mind that cytokine production elicited by TLR-activation can also occur involving NF-κB-independent pathways.

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Declaration of interests
There are no conflicts of interests with this work.
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