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To cite this version:

R. Vrancken, A. Haegeman, J. Dewulf, J. Paeshuyse, G. Puerstinger, et al.. The reduction of CSFV transmission to untreated pigs by the pestivirus inhibitor BPIP: a proof of concept. Veterinary Microbiology, Elsevier, 2009, 139 (3-4), pp.365. <10.1016/j.vetmic.2009.06.026>. <hal-00526939>

HAL Id: hal-00526939
https://hal.archives-ouvertes.fr/hal-00526939
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Accepted Manuscript

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PII: S0378-1135(09)00311-3
DOI: doi:10.1016/j.vetmic.2009.06.026
Reference: VETMIC 4481

To appear in: VETMIC

Received date: 1-12-2008
Revised date: 3-6-2009
Accepted date: 12-6-2009

Please cite this article as: Vrancken, R., Haegeman, A., Dewulf, J., Paeshuyse, J., Puerstinger, G., Tignon, M., Le Potier, M.-F., Neyts, J., Koenen, F., The reduction of CSFV transmission to untreated pigs by the pestivirus inhibitor BPIP: a proof of concept, Veterinary Microbiology (2008), doi:10.1016/j.vetmic.2009.06.026

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The reduction of CSFV transmission to untreated pigs by the pestivirus inhibitor BPIP: a proof of concept

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Keywords: Antivirals, pestivirus, classical swine fever virus, virus transmission

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Abstract.

5-[(4-bromophenyl)methyl]-2-phenyl-5H-imidazo[4,5-c]pyridine (BPIP) is a representative molecule of a novel class of highly active *in vitro* inhibitors of the replication of Classical swine fever virus (CSFV). We recently demonstrated in a proof of concept study that the molecule has a marked effect on viral replication in CSFV-infected pigs. Here, the effect of antiviral treatment on virus transmission to untreated sentinel pigs was studied. Therefore, BPIP-treated pigs (n=4), intramuscularly infected with CSFV, were placed into contact with untreated sentinel pigs (n=4). Efficient transmission of CSFV from four untreated seeder pigs to four untreated sentinels was observed. In contrast, only two out of four sentinel animals in contact with BPIP-treated seeder animals developed a short transient infection, of which one was likely the result of sentinel to sentinel transmission. A significant lower viral genome load was measured in tonsils of sentinels in contact with BPIP treated seeder animals compared to the positive control group (p=0.015). Although no significant difference (p=0.126) in the time of onset of viraemia could be detected between the groups of contact animals, a tendency towards the reduction of virus transmission was observed. Since sentinel animals were left untreated in this exploratory trial, the study can be regarded as a worst case scenario and gives therefore an underestimation of the potential efficacy of the activity of BPIP on virus transmission.
Classical swine fever (CSFV), Bovine viral diarrhea virus (BVDV) and Border
disease virus (BDV) belong to the family of Flaviviridae, genus pestivirus (van
Regenmortel et al., 2000) and can, in case of an outbreak, result in major economical
consequences for the affected region (Domenech et al., 2006). In particular CSFV has
been responsible for major economic losses (Greiser-Wilke et al., 2007). Currently,
CSF outbreaks are controlled by a stamping-out policy and pre-emptive eradication of
neighbouring herds. These measures proved efficient, but the slaughter of large
numbers of often healthy, uninfected pigs is increasingly criticized by the public
opinion (van Oirschot, 2003; Le Potier et al., 2006).

As an alternative/additional control strategy, the use of an antiviral treatment for the
containment of outbreaks of infectious diseases of livestock like foot-and-mouth
disease and CSF has been proposed (Goris et al., 2008; Vrancken et al., 2008). For
CSFV, we previously reported on the in vitro inhibition of viral replication by BPIP, a
representative of a new class of imidazopyridines, specifically targeting the viral
RNA-dependent RNA-polymerase (Vrancken et al., 2008). Subsequent in vivo studies
revealed that BPIP was able to significantly reduce the viral load in CSF-infected pigs
(Vrancken et al., 2009). It was the purpose of this study to assess the effect of BPIP
treatment on the transmission of CSFV from infected pigs to untreated, naive pigs.

An animal experiment was designed, according to the French legislation on animal
experimentation, and carried out at the high containment facilities at the Agence
Française de Sécurité Sanitaire des Aliments (AFSSA Ploufragan, France). Groups of
four, twenty-nine-week old, Specific Pathogen Free (SPF) Large White pigs (ca. 30
kg) of mixed sex, originating from protected breeding facilities at AFSSA Ploufragan,
were held in isolation units. One group of four SPF pigs received BPIP containing
feed at 75 mg/kg/day for a period of 15 consecutive days. BPIP synthesis and formulation was carried out as described earlier (Puerstinger et al., 2006; Vrancken et al., 2009). A positive and negative control group was held in a separate isolation unit and received normal feed. One day after the first administration of BPIP, the animals of the BPIP-treated and positive control group were infected intramuscularly with 4 ml of $10^{4.5}$ TCID$_{50}$/ml of the CSFV field-isolate Wingene (subgroup 2.3, Vanderhallen et al., 1999). Two days post infection (dpi) four untreated sentinel animals were placed in an adjacent pen allowing nose to nose contact between infected and sentinel pigs.

All experimentally infected animals (BPIP-treated and untreated seeder pigs) were clinically observed on a daily basis until 33 dpi and all sentinel animals until 40 dpi. During this observation period all animals were blood sampled three (0-18 dpi) or two (19- 40 dpi) times a week. After the observation period, all animals were euthanized and tonsils were sampled. All blood and organ samples were analyzed by means of virus isolation (VI) and real-time RT-PCR (TaqVet PPC, LSI, France) as described earlier (Vrancken et al., 2009). The real-time RT-PCR assay, using $\beta$-actin as an internal control, had a limit of detection of $2.2 \pm 1.2$ equivalent genome copies (EGC). Samples with a positive signal but a viral load below $2.2$ EGC for a 5 µl reaction were considered as not quantifiable.

As presented in Fig. 1A, BPIP-treatment had a marked effect on the period of viraemia; three out of four BPIP-treated seeder pigs developed a short transient vireamia of which two animals were only positive for 2 days and one animal for 7 days. The remaining animal tested negative in VI during the whole period of observation. In contrast, the untreated seeder pigs (positive control group) tested invariably positive from 5 dpi until death (22 dpi) or 26 dpi. One animal scored
positive until the end of the experiment (33 dpi) (Fig. 1B). Analysis of the blood samples of the sentinel pigs in contact with BPIP-treated seeder pigs revealed one positive animal at 14 dpi and a second as late as 26 dpi. The two remaining animals remained negative during the whole period of observation (Fig. 1A). All sentinel pigs in contact with the untreated seeder group became viraemic on VI between 14 and 19 dpi. Three out of four animals remained VI positive in blood during the whole period of observation and one contact pig of this group showed a transient infection between 19 and 26 dpi (Fig 1B). The average time until onset of viraemia (VI) in the contact groups was calculated and although a tendency towards later infection could be observed in the sentinel pigs in contact with the BPIP-treated seeder group compared to those in contact with the untreated seeder group (20.00 ± 8.49 days vs. 15.75 ± 2.36 days), this difference was not statistically significant (p=0.126 [Cox regression survival analysis]).

The effect of a BPIP-treatment on viraemia was substantiated by real time RT-PCR analysis where in the BPIP-treated seeder group one animal with a very low level of CSFV-genome was detected at 2 dpi (EGC not quantifiable). Between day 5 and 16 post experimental inoculation, all BPIP-treated seeder animals scored positive (log$_{10}$EGC between 0.55 ± 0.15 and 2.52 ± 0.8). At 16 dpi, three out of four animals tested positive with an average log$_{10}$EGC of 0.69 ± 0.32. In all four untreated seeder pigs the presence of viral genome was detected throughout the observation period from day 2 post infection until death (22 dpi) or the end of the experiment (33 dpi) with log$_{10}$EGC between 0.46 ± 0.28 and 6.03 ± 0.98.

As depicted in Fig. 2, real-time RT-PCR results revealed a significant lower viral genome load in the sentinel group in contact with BPIP-treated seeder pigs than the group in contact with the untreated seeder pigs. In one out of four animals in contact
with BPIP-treated seeder animals detectable levels of viral RNA were observed from 14 dpi until the end of the study (40 dpi). A second animal became positive between days 26 and 40 post inoculation. As late as 40 dpi, a very low level of viral RNA (log_{10} EGC=0.81) was detected in a third animal and one animal remained negative during the whole observation period. In contrast, the group of sentinel pigs in contact with the untreated seeder pigs, viral RNA could be detected in two out of four animals at 14 dpi. A further animal tested positive at 16 dpi and a fourth animal at 19 dpi. All animals remained positive until the end of the experiment (40 dpi).

At the end of the experiment, infectious virus could be isolated from the tonsils of one animal in contact with BPIP-treated seeder pigs. Three out of four animals in contact with untreated seeder pigs scored positive in VI (Table 1). Real-time RT-PCR analysis revealed that three out of four pigs in contact with BPIP-treated seeder pigs showed a significantly lower viral genome load in the tonsils compared to the sentinel pigs in contact with the untreated seeder animals (p=0.015 [Two tailed students t-test, unequal variances]) (Table 1).

The VI and real-time RT-PCR results revealed that only one animal developed a short transient infection with an onset comparable to the positive control group (14 dpi), which may be explained by the fact that the BPIP-treated seeder pigs had only a very short period of viraemia. The source of infection of the second VI-positive animal could not be determined in this experimental set-up. A direct infection by contact with the BPIP-treated seeder animals is however unlikely considering the fact that no infectious virus could be isolated from these animals after day 12 pi and that infection of the sentinel could only be demonstrated on VI as late as 26 dpi. Furthermore, because the time interval between the initially infected sentinel pig and the second sentinel pig (12 days), it is reasonable to assume that this infection did not originate
from the seeder pigs, but was most likely due to a secondary transmission from the initially infected contact pig (Ribbens et al., 2004). The same conclusion can likely be drawn for the third animal that, although no infectious virus could be isolated, scored positive on real-time RT-PCR as late as 40 dpi. Although real-time RT-PCR results indicate that a BPIP-treatment resulted in a significant lesser virus transmission towards untreated pigs, it did not result in the observation of a significant later onset of viraemia in the untreated animals in contact with the BPIP-treated seeder group, due to the limited power of the experiment (only 4 contact pigs in each group). Additional experiments must therefore be performed to confirm the here observed tendency towards reduction of virus transmission and to determine the statistical significance.

This exploratory trial can be regarded as a worst-case scenario where both treated and untreated animals were in close contact. Since the aim of an antiviral treatment is not to protect individual animals, but to curb viral spread between herds, all animals within a herd would be treated and therefore the currently obtained results are probably an underestimation of the potential efficacy of an antiviral treatment to reduce virus transmission. Furthermore, both the previously published study of the effect of BPIP on CSFV-viraemia (Vrancken et al., 2009), and the current, were carried out with a lead molecule of this class. Further optimization of the antiviral efficacy could lead to further decrease (if not complete suppression) of the period of viraemia and consequently on virus transmission.

In conclusion, our preliminary findings indicate a reduction of virus transmission from CSFV-infected BPIP-treated animals to untreated sentinel pigs and further trials with BPIP (or a more potent analogue) will be performed to further confirm the observed trend.
Aknowledgements.

This work was supported by a grant (S6146 section 2) from the research foundation DG4 of the Belgian Government (“Federale Overheidsdienst Volksgezondheid, Veiligheid van de Voedselketen en Leefmilieu”) and by the EU Network of Excellence, EPIZONE (Contract No FOOD-CT-2006-016236). J. Paeshuyse is a post-doctoral fellow of the “Fonds voor Wetenschappelijk Onderzoek, Vlaanderen”. We thank Cariolet R., Rault J.-C. and Hutet E. for their assistance during the animal trial and Debaugnies R., Denne M., Jebbari F., and Thoraval C. for their technical assistance during this study.

References.


Tables

Table 1. Virus Isolation (VI) and real-time RT-PCR (expressed as log$_{10}$(EGC)) results of tonsils of untreated sentinel pigs in contact with BPIP-treated (S1 → S4) and untreated (PS1 → PS4) animals.

<table>
<thead>
<tr>
<th>Animal</th>
<th>VI</th>
<th>log$_{10}$(EGC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>+</td>
<td>3.89</td>
</tr>
<tr>
<td>S2</td>
<td>-</td>
<td>3.89</td>
</tr>
<tr>
<td>S3</td>
<td>-</td>
<td>neg</td>
</tr>
<tr>
<td>S4</td>
<td>-</td>
<td>3.89</td>
</tr>
<tr>
<td>Average:</td>
<td></td>
<td>3.89</td>
</tr>
<tr>
<td>PS1</td>
<td>+</td>
<td>7.66</td>
</tr>
<tr>
<td>PS2</td>
<td>-</td>
<td>5.26</td>
</tr>
<tr>
<td>PS3</td>
<td>+</td>
<td>7.31</td>
</tr>
<tr>
<td>PS4</td>
<td>+</td>
<td>6.29</td>
</tr>
<tr>
<td>Average:</td>
<td></td>
<td>6.63 ± 1.08</td>
</tr>
</tbody>
</table>

Viral genome loads of animals in contact with BPIP-treated pigs are significantly lower compared to animals in contact with the positive control group (p=0.015).
Legends to the figures

Fig. 1. Duration of viraemia in blood after infection with CSFV Wingene and transmission to untreated sentinel pigs from A) BPIP-treated/Infected (1 → 4) to untreated sentinel (S1 → S4) animals; B) Untreated/Infected (P1 → P4) to untreated sentinel (PS1 → PS4) animals. Ⓐ: animal euthanized at 22 days post infection; BPIP-treated and untreated seeder animals were euthanized at 33 days post infection, contact sentinel pigs at 40 days post infection.

Fig. 2. Mean virus genome load in blood of untreated sentinel animals in contact with BPIP-treated (diamonds; n=4) and untreated (squares; n=4) seeder pigs as determined by means of real-time RT-PCR. Number of animals tested positive at a given time point is shown between brackets. Statistical significance (p-values) is given at each measureable timepoint.
Figure 1

Virus Isolation in Blood

Days post experimental inoculation

Virus Isolation in Blood

Days post experimental inoculation
Figure 2

Days post experimental infection

Equivalent Genome Copies (log_{10}-transformed)