Contreras B, G. Andres; Guterbock, Walter M.; Muñoz R, Juan; Sears, Phillip M.
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Universidad de Córdoba
Montería, Colombia

Available in: http://www.redalyc.org/articulo.oa?id=69325829012
Comparison of systemic and intramammary dry cow treatments

Comparación de terapias sistémicas e intramamarias para el periodo seco en vacas lecheras

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ABSTRACT

Objective. To compare four different dry cow treatments (DCT) and establish their effectiveness in reducing intramammary infections (IIM). Materials and methods. DCTs included systemic tylosin (12g) alone or accompanied by cefapirine intramammary infusions and or an internal teat sealant. A total number of 278 cows at the end of lactation period were randomly assigned to one of 4 dry cow treatment groups: CESE Group (n=89), intramammary cephapirin and teat sealant. TYCESE Group (n=84), intramammary cephapirin, tylosin 12 g intramuscular and teat sealant. TYSE Group (n=86), 12 g intramuscular tylosin and teat sealant; TY Group (n=76) 12 g intramuscular tylosin only. Milk samples for culture were collected at dry-off and 1 and 2 weeks after calving. Somatic cell counts (SCC) were taken from Dairy Herd Improvement Association (DHI) tests at dry-off, and the first two test days after calving. Results. Bacteria cure rate for Gram-positive intramammary infections (IMI) for TYCESE group was 93.6%, CESE group 78.9%, TYSE group 88.2%, and TY group 78.1%. All four groups showed a decrease in the SCC upon the first and second test after calving. Conclusions. The use of systemic tylosin in combination with intramammary antibiotics increased DCT effectiveness improving the Gram-positive cure rate IMI. Furthermore, systemic tylosin alone plus teat sealant is as effective as cephapirin plus teat sealant when used as DCT.

Key words: Dry period, mastitis, tylosin (Source: CAB).

RESUMEN

Objetivo. Comparar 4 tratamientos para el periodo seco (TPS) y establecer su efectividad en reducir infecciones intramamarias (IIM). Materiales y métodos. Los TPS incluían tilosin sistémica (12 g) solamente o acompañada con infusions intramamarias de cefapirina y/o un sellante interno de pezones. Un total de 278 vacas al final de la lactancia fueron asignadas al azar a 1 de 4 grupos de tratamiento. Grupo CESE (n=89), cefapirina y sellante interno de pezones. Grupo TYCESE (n=84), cefapirina intramamaria, tilosina intramuscular y sellador interno de pezones. Grupo TYSE (n=86), tilosina intramuscular y sellante interno de pezones. Grupo TY (n=76) tilosina intramuscular solamente.
Muestras de leche de cada cuarto productivo fueron tomadas al momento del secado, y 1 y 2 semanas después del parto. Los recuentos individuales de células somáticas (RCS) fueron tomados del control lechero (DHI) en sus pruebas de antes del momento de secado, y las primeras dos pruebas después del parto. **Resultados.** Las tasa de cura bacteriológica para IIM causadas por Gram positivos en TYCESE fue de 93.6%, para CESE 78.9%, TYSE 88.2% y TY 78.1%. Todos los grupos de tratamiento mostraron una disminución en RCS a la primera y segunda prueba después del parto en comparación con la de secado. **Conclusiones.** El uso de tilosina en combinación con cefapirina incremento la efectividad del TPS, mejorando las tasas de cura de IIM por Gram positivos. Además TYSE fue tan efectivo como CESE en la reducción de IIM.

**Palabras Clave:** Mastitis, periodo seco, tilosina (Fuente: CAB).

INTRODUCTION

Dry cow therapy (DCT) is defined as the use of antibiotics immediately after the last milking of lactation. There are three different approaches to DCT. First, intramammary dry treatment alone is widely used and very effective in reducing intramammary infections (IMI) (1, 2). However, low efficacy against some coagulase negative staphylococci (CNS) IMI has been demonstrated. (3) Another approach is systemic dry cow therapy, which is relatively inexpensive and easy to use but has been reported to be less effective.

For example, Nickerson et al (4) administered tilmicosin subcutaneously and found it to be less effective against IMI caused by *Staphylococcus aureus* than intramammary infusions of cephapirin or tilmicosin. Finally a combination of systemic and intramammary antibiotics can also be administered. However, its efficacy depends upon the antibiotic class pharmacological characteristics. In fact, Erskine et al (5) reported a low efficacy against *Staphylococcus aureus* IMI when using oxytetracycline hydrochloride systemically in combination with intramammary cephapirin benzathine. The low efficacy of this treatment could have been related to the poor diffusion of oxytetracycline into the mammary gland.

The use of tylosin as systemic DCT was reported to be as effective as intramammary infusion with a preparation with beta lactams and streptomycin against CNS (6). However, combined DCT including systemic tylosin has not been previously reported. This antibiotic belongs to the macrolide family of antibiotics. One of the main advantages of this type of antimicrobials is an excellent diffusion into the mammary gland related to its basic pK (pH at which concentrations of dissociated and undissociated antibiotic are equal), which results in a very high milk to plasma concentration ratio of 5:1 (7,8). Once the macrolides are in milk they are virtually trapped inside the mammary gland. The objective of this study was to compare four different DCTs and establish the effectiveness of systemic (intramuscular) tylosin therapy as DCT at quarter level when was added to intramammary infusion of cephapirin, a teat sealant, or alone as systemic DCT.

MATERIALS AND METHODS

**Study site.** Cows on a large commercial farm in Michigan were selected. Every week, for two months, and after confirming their gestation to be greater than 150 days, a group of Holstein cows in their second, third or fourth lactation due to go dry were assigned to one of four DCT groups. A completely randomized design that included three treatments and a control was used (Table 1).

Table 1. Summary of gram-positive intramammary infections by treatment groups at cow and quarter level.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>CESE</th>
<th>TYCESE</th>
<th>TYSE</th>
<th>TY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cows Enrolled</td>
<td>75</td>
<td>71</td>
<td>72</td>
<td>60</td>
</tr>
<tr>
<td>Cows with IMI at dry-off (% cows infected)</td>
<td>30</td>
<td>28</td>
<td>33</td>
<td>32</td>
</tr>
<tr>
<td>Cow cured after DCT (% cows cured)</td>
<td>20</td>
<td>21</td>
<td>17</td>
<td>15</td>
</tr>
<tr>
<td>Cows with new IMI after calving</td>
<td>4</td>
<td>4</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>Quarters Enrolled</td>
<td>289</td>
<td>271</td>
<td>276</td>
<td>226</td>
</tr>
<tr>
<td>Non-functional quarters</td>
<td>11</td>
<td>13</td>
<td>12</td>
<td>14</td>
</tr>
<tr>
<td>Quarters infected at dry-off (% quarters infected)</td>
<td>57</td>
<td>47</td>
<td>68</td>
<td>73</td>
</tr>
<tr>
<td>Quarters cured after DCT (% quarters cured) (78.9%)</td>
<td>45</td>
<td>44</td>
<td>60</td>
<td>57</td>
</tr>
</tbody>
</table>

1Treatment designations: CESE cephaprin 300 mg intramammary and Orbeseal® (4 g of internal teat sealant containing 65% bismuth subnitrate); TYCESE: tylosin 12 g intramuscular; cephaprin 300 mg intramammary, and Orbeseal®; TYSE: Tylosin 12 g intramuscular and Orbeseal®; TY: tylosin 12 g intramuscular. *Different superscripts within each row differ (p<0.05).
**Treatments.** Treatment CESE (n=89 cows), reflecting standard practice on the farm, was used as control. After the last milking of lactation, animals were first infused in each productive quarter with a commercial preparation of 300 mg of cephapirin (Tomorrow®, Fort Dodge, Ft Dodge, Iowa, USA) intramammarly and then with 4 g of commercially available internal teat sealant containing 65% bismuth subnitrate (Orbeseal®, Pfizer. New York, New York, USA). Treatment TYCESE (n=84 cows) animals were first infused in each productive quarter with cephapirin intramammary 300 mg, then injected intramuscularly with tylosin (Tylosin, Agripharma. Westlake, Texas, USA) 12 g and finally were infused with teat sealant. Group TYSE (n=86 cows) animals received tylosin 12 g intramuscular and then teat sealant was infused in each productive quarter. Group TY (n=76 cows) animals received 12 g tylosin intramuscularly. Intramammary infusions were administered by the partial insertion technique.

**Prior to instillation.** All teat ends were prepared following the farm’s milking preparation routine that included a predip solution containing 0.1% iodine, after which the dry teat end was disinfected with a cotton pad containing 70% isopropyl alcohol. Intramuscular injections were given in the neck, splitting the dose between two different sites. Tylosin dosage calculation was based in 20 mg/Kg dose for a 600 Kg cow (average weight for mature cows on this commercial farm).

**Milk samples.** Quarter milk samples were taken following National Mastitis Council guidelines (9) from all functional quarters at dry-off and at 1 and 2 wks after calving. Samples were immediately processed at the dairy’s on-farm laboratory. A 10 µL aliquot from each sample was cultured on blood agar containing 0.5% esculin at 37°C for 24 h. Colonies were tentatively identified as CNS, streptococci, *Staphylococcus aureus*, coliforms or others; a presumptive diagnosis of CNS, streptococci, coliform, or other pathogens was made, based on colony growth, morphology and appearance, pattern of hemolysis, and catalase reaction. *Staphylococcal* isolates were tested for coagulate production with the tube coagulate test. Gram-negative bacteria were plated on MacConkey agar to facilitate identification. Only CNS, streptococci and *S. aureus* were included in data analysis. This approach was based on the label use for cephapirin (Tomorrow®) and Tylosin sold in the United States at the time of the trial, not on their antibacterial activity. A cow was considered infected if at least one productive quarter had an IMI. A quarter was considered infected if five or more cfu of the same kind were identified, and was considered contaminated if three or more different colony types were present. Post calving samples were used to establish bacterial cure. A cow was considered cured if she had an IMI at dry-off and all productive quarters were negative at both samples post-calving. A cow was not cured if she had an IMI at dry-off and continued with the same quarter(s) infected with the same species at first and second test after calving. A quarter was considered cured if it was positive at dry-off and negative in first and second post-calving samples. It was considered not cured if it was positive to the species found at dry-off and again 1 and/or 2 wk after calving. A quarter was considered newly infected if it was negative at dry-off and positive at 1 and 2 wk after calving Dairy Herd Improvement (DHI) tests were recorded to monitor SCC.

The last test of lactation was considered the dry-test, and then linear SCC scores from subsequent monthly tests, were used to determine changes in SCC. Milk production records were collected by DHI personnel and total production was established as projected to 305 days mature equivalent (ME) by a herd management software (DairyComp®, Herd Management software, Tulare, California, USA) based on milk production at DHI test taken between 180 and 200 days in milk (DIM) after calving following the DCT assigned. A composite milk sample was taken at calving to perform antibiotic residue test (Delvotest® system, DMS food specialties, Parsippany, New Jersey, USA). Cows with less than 30 d dry were excluded because of antimicrobial residue issues that required milk withdrawal. Cows with more than 100 d dry were also excluded. Of the 335 cows enrolled in the dry cow treatment trial, 278 had complete records and were included in the results.

**Analysis of results.** Culture results were analyzed with the PROC GLIMMIX procedure for binary variables (SAS Inst. Inc., Cary, North Carolina, USA). Results were blocked by sampling date and the random statement was assigned to quarter within cow as experimental unit. The SCC scores were analyzed as repeated measures using a mixed model procedure (PROC MIXED; SAS Inst. Inc., Cary, North Carolina, USA) following the equation:

\[
Y_{ijk} = \mu + G_i + I_j + G_i \times I_j + S_k + S_k \times G_i + S_k \times I_j + S_k \times G_i \times I_j + E_{ijk}
\]

Where \(Y_{ijk}\) is the dependent variable SCC score for a cow in group \(i\), with infection status \(j\), at
sample k as repeated measure, and $E_{ijk}$ is the random error assumed to be correlated. The Satterthwaite's method for estimating degrees of freedom was used. Significance level was set at $\alpha=0.05$.

**RESULTS**

A total of 123 cows (44%) had IMI at dry-off and no differences in infection rate were found when comparing all four groups. Percentages of cows with IMI within each treatment group were as follows CESE 40%, TYCESE 39%, TYSE 45% and TY 53%. Cure rates at cow level were significantly higher for group TYCESE (75%) when compared to group TY (46%). Group CESE had a (66%) cure rate and TYSE (51%) and this difference was not significant. A summary of these results is shown in Table 1. Despite of the use of antibiotics, residue tests were negative on all samples taken after calving.

A total of 245 quarters (23% overall) were found infected with CNS, streptococci or *S. aureus*. IMI rates by quarters at dry-off are shown in table 1. Group TY had 32.3% infected quarters and group TYSE had 24.6%, both were significantly higher ($p<0.05$) than TYCESE 17.3% and CESE 19.7%. Gram-positive bacterial infection results are summarized in table 2.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNS% quarters infected at dry-off (% quarters infected)</td>
<td>41(72%)a</td>
<td>27(57%)b</td>
<td>28(41%)b</td>
<td>58(79%)a</td>
</tr>
<tr>
<td>CNS% quarters cured after DCT</td>
<td>73%a</td>
<td>92%b</td>
<td>89%b</td>
<td>72%a</td>
</tr>
<tr>
<td>Streptococci quarters infected at dry-off (% quarters infected)</td>
<td>14(24%)</td>
<td>18(38%)</td>
<td>34(50%)</td>
<td>14(19%)</td>
</tr>
<tr>
<td>Streptococci quarters cured after DCT</td>
<td>100%</td>
<td>94%</td>
<td>91%</td>
<td>100%</td>
</tr>
<tr>
<td><em>S. aureus</em> quarters infected at dry-off</td>
<td>2</td>
<td>2</td>
<td>6</td>
<td>1</td>
</tr>
</tbody>
</table>

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CNS accounted for 62.9% of the infections, streptococci for 32.6% and *S. aureus* for 4.5%. The staphylococcal infection rate was significantly higher in group TY when compared to TYCESE and TYSE ($p<0.05$) (Table 1). Infections caused by *S. aureus* were found in 2 quarters of group CESE, 2 quarters of TYCESE, 6 quarters of TYSE and 1 quarter of TY. With such a small number of infections it was not possible to establish differences among groups. Furthermore, IMI with *S. aureus* exhibit intermittent shedding that requires at least 3 multiple samplings to identify a truly infected quarter and this trial only collected 1 sample at dry-off (10). Further studies are needed to evaluate tylosin efficacy alone or in combination with other antibiotics and or teat sealants against *S. aureus* IMI. Coliform infections were found in 5 quarters from 3 different cows at the moment of dry-off (results not shown). These infections were cleared after parturition.

Bacterial cure rates for gram-positive bacteria by quarters are shown in table 2. TYCESE had the highest cure rate (93.6%) and was significantly different ($p<0.05$) from TY (78.1%) and CESE (78.9%). No difference was found when TYCESE was compared to Group TYSE (88.2%). New infections rates were low, group CESE (1.8%), group TYCESE (2.6%), group TYSE (2.9%) and group TY (3.3%). When comparing cure rates by bacteria type (Table 1) all four treatments had good efficacy against streptococcal infections, but there were differences in cure rates for staphylococcal infections. Group TYCESE had higher cure rates (92.5%) than CESE (73.17%) and TY (72.4%) ($p<0.05$).

All four treatments resulted in a decrease in the SCC at the first and second test after calving and no statistical differences were observed among them (Figure 1). Milk production at test date between 180 to 200 DIM was analyzed using 305 ME value. Milk production for each treatment group was as follows: TYCESE 23.910±5.686 lb, CESE 22.056±5.104 lb, TY 21.900±4.278 lb and TYSE 21.791±4.086 lb. When the difference between arithmetic mean of previous and current lactations was calculated for all groups, no significant differences were found.

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**Figure 1.** logSCC at dry-off, first test and second test after calving for each treatment group. Values are means ± SEM.
DISCUSSION

Bacterial cure rates for systemic tylosin (group TY 78.1%) are similar to those obtained by McDougall et al (11) in New Zealand, who administered tylosin intramuscularly to treat clinical mastitis during lactation and obtained 82% cure rates for Gram-positive infections. Systemic tylosin as DCT alone (TY) or plus teat sealant (TYSE) is as effective as intramammary cepapirin plus teat sealant (Group CESE 78.9%) to reduce IMI.

The randomization was effective in distributing IMI among groups at cow level, but was ineffective at quarter level generating a bias, observed in a higher number of quarter infections and a lower cure rate for the TY group. An ideal randomization would have included blocking after culturing quarters but this process would have represented major logistical problems on this commercial farm. However, all four DCTs were effective in reducing IMI.

Nickerson et al (4) used a systemic macrolide (tylmicosin) to treat IMI during the dry period reporting low cure rates against staphylococcal infections, however, the dose (1 mg/Kg) used by the authors does not have bactericidal activity, and subcutaneous administration could have delayed antibiotic distribution. Tylosin’s effectiveness during this trial could be related to its distribution properties and the fact that a higher than recommended dose with a single injection was used. Tylosin’s peak milk concentration was reported to be 10 µg/ml after a single injection at a dose of 20 mg/Kg b.w. (12) and 18 µg/ml after three repeated injections at a dose of 10 mg/Kg b.w (13).

In the present work, tylosin’s dose was based on average calving weight for cows (600 Kg) at 20 mg/Kg. Despite of the relative high dose used, antibiotic residues were not found in the composite samples taken at calving. It is presumable that peak milk concentration in tylosin treated animals was higher than previously reported, therefore achieving bactericidal activity, which was observed for macrolides when used in high concentrations (14).

When analyzing cure rate by bacteria type, intramammary treatment alone could have been limited in its ability to eliminate infections during the dry period (CESE 73.2%). Similar to what was reported by Bolurchi et al (6) inclusion of systemic tylosin in DCT increased cure rates for CNS IMI. Because CNS could be responsible of nearly half of the IMI during the dry period, an increase in the cure rate towards this bacteria type obtained by the addition of systemic tylosin to the intramammary DCT would have an important economic impact (15).

In conclusion, DCT remains a cost-effective measure compared with no treatment at dry-off (3,16,17). The use of systemic tylosin in combination with the intramammary cephapirin increased the effectiveness of intramammary DCT against Gram-positive IMI. Though, inclusion of a combined systemic and intramammary treatment has to be based on an economic analysis of the increased cost of such therapy versus its higher effectiveness. Interestingly, there were no differences between the use of tylosin plus teat sealant and intramammary cephapirin plus teat sealant at dry-off. Therefore the use of these 2 DCT options may be based on the logistical advantages of each treatment protocol. Tylosin may have the advantage of being easier to administer in dairies with confinement housing where headlocks are available to treat large groups of animals. Finally, it is important to mention that adding the teat sealant to the systemic treatment with tylosin at dry-off improved the response of the treatment. Further studies are needed to validate this effect in other dairy herds and to test the use of extended therapy with tylosin particularly when selective DCT is used.

REFERENCES


