ABSTRACT

Our objective was to evaluate time to pregnancy after the first service postpartum and pregnancy per artificial insemination (P/AI) in dairy cows managed with 2 resynchronization of ovulation programs. After first service, lactating Holstein cows were blocked by parity (primiparous vs. multiparous) and randomly assigned to the d 32 Resynch (R32; n = 1,010) or short Resynch (SR; n = 1,000) treatments. Nonpregnancy diagnosis (NPD) was conducted 32 ± 3 d after AI by transrectal ultrasonography. Nonpregnant cows in R32 received the Ovsynch protocol: GnRH, PGF2α 7 d later, GnRH 56 h later, and timed AI (TAI) 16 to 18 h later. Cows in SR with a corpus luteum (CL) ≥15 mm and a follicle ≥10 mm at NPD received PGF2α, PGF2α 24 h later, GnRH 32 h later, and TAI 16 to 18 h later. Cows in SR without a CL ≥15 mm or a follicle ≥10 mm at NPD received a modified Ovsynch protocol with 2 PGF2α treatments and progesterone (P4) supplementation (GnRH plus CIDR, PGF2α and CIDR removal 7 d later, PGF2α 24 h later, GnRH 32 h later, and TAI 16 to 18 h later). Blood samples were collected from a subgroup of cows at the GnRH before TAI (R32 = 114; SR = 121) to measure P4 concentration. Binomial outcomes were analyzed with logistic regression and hazard of pregnancy (R32 = 485; SR = 462) with Cox’s proportional regression in SAS (SAS Institute, Cary, NC). For P/AI analysis, the TAI service was the experimental unit (R32 = 720; SR = 819). Models included treatment and parity as fixed effects and farm as random effect. The hazard of pregnancy was greater for the SR treatment (hazard ratio = 1.18; 95% confidence interval: 1.01–1.37). Median time to pregnancy was 95 and 79 d for the R32 and SR treatments, respectively. Treatment did not affect overall P/AI 32 ± 3 d after AI (R32 = 31.0% vs. SR = 33.9%) or for cows with a CL at NPD (R32 = 32.7% vs. SR = 32.8%). For cows with no CL at NPD, P/AI was greater for the SR treatment (36.9%) than for the R32 treatment (28.6%). Pregnancy loss from 32 to 63 d after AI was similar for all services combined (R32 = 8.3% vs. SR = 10.4%) and for cows with no CL at NPD (R32 = 13.2% vs. SR = 7.2%) but tended to be affected by treatment for cows with a CL at NPD (R32 = 6.8% vs. SR = 11.9%). Treatment affected the proportion of cows with P4 ≤0.5 ng/mL at the GnRH before TAI for all cows (R32 = 70.0% vs. SR = 81.8%), tended to have an effect among cows with a CL (R32 = 70.0% vs. SR = 81.8%), and had no effect for cows with no CL (R32 = 64.7% vs. SR = 81.8%). We concluded that the SR program reduced time to pregnancy because of a reduction of the interbreeding interval for cows with a CL at NPD and greater P/AI in cows with no CL at NPD.

Key words: resynchronization, corpus luteum, dairy cow, timed artificial insemination

INTRODUCTION

Dairy farm profitability depends on the reproductive performance of cows, which is primarily determined by the rate at which they become pregnant after the end of the voluntary waiting period. Despite recent gains on first-service pregnancy per AI (P/AI) due to improved dairy herd management (Wiltbank and Pursley, 2014), a substantial proportion of cows need immediate reinsemination after the first service. Therefore, to minimize the interbreeding interval, many farms use reproductive management strategies for second and greater AI services that combine insemination at detected estrus and timed AI (TAI) after resynchronization of ovulation with Ovsynch-type protocols (Pursley et al., 1995; commonly referred to as Resynch). These protocols are usually initiated at the time of or 7 d before nonpregnancy diagnosis (NPD; Fricke et al., 2003; Bartolome et al., 2005; Giordano et al., 2012c). Farms that combine AI at detected estrus and TAI can initiate Resynch as early as 25 ± 3 d after AI. Although
this strategy can be beneficial because it shortens the interbreeding interval for cows that receive TAI, the first GnRH treatment of the protocol coincides with the time at which many cows are expected to display estrus (Remnant et al., 2015; Wijma et al., 2017). The GnRH-induced LH surge reduces estrus expression (Mendonça et al., 2012; Bruno et al., 2014; Wijma et al., 2017) through induction of ovulation or by suppressing the estradiol surge responsible for estrus behavior (Jo and Fortune, 2003). Further, the pregnancy status of cows at the time of the GnRH treatment 25 ± 3 d after AI is unknown. Therefore, many pregnant cows receive an unnecessary treatment, thus increasing treatment costs and disrupting cow normal behavior.

To take advantage of a short interbreeding interval for TAI services while avoiding a reduction in the proportion of cows undergoing AI at detected estrus before NPD and unnecessary treatment of pregnant cows with GnRH, we evaluated a reproductive management strategy based on the ovarian structures present at NPD 32 ± 3 d after AI (Wijma et al., 2017). Cows with a corpus luteum (CL) ≥15 mm and a follicle ≥10 mm at NPD (hereafter referred to as CL cows) received a resynchronization of ovulation protocol without an initial GnRH treatment to induce a new follicular wave (short Resynch; PGF$_{2\alpha}$, PGF$_{2\alpha}$ 1 d later, GnRH 32 h later, and TAI 16 to 18 h after GnRH). On the other hand, cows not expected to respond to the short Resynch protocol based on their ovarian status (i.e., no CL ≥15 mm or no follicle ≥10 mm; hereafter referred to as no-CL cows) received a modified Ovsynch protocol (i.e., 2 PGF$_{2\alpha}$ treatments 24 h apart) with progesterone (P$_4$) supplementation. This management strategy was compared with a similar program in which all cows received GnRH treatment 7 d before NPD. Removing the GnRH treatment 25 ± 3 d after AI resulted in approximately 17% more cows inseminated at detected estrus, but it also resulted in a P/AI reduction of approximately 8 percentage points for TAI services in CL cows. Nevertheless, the cumulative proportion of cows pregnant after AI at detected estrus and TAI was similar because of the greater number of pregnancies generated through insemination of cows at detected estrus before NPD in the short Resynch treatment. Although results from this experiment (Wijma et al., 2017) were promising, additional research is needed to determine whether the strategy based on ovarian status at NPD is superior to traditional programs combining AI at detected estrus and TAI after blanket use of resynchronization of ovulation. Moreover, it is necessary to corroborate that removal of the initial GnRH treatment does not compromise P/AI for CL cows to an extent that may offset the benefit of shorter interbreeding interval.

Thus, we hypothesized that a resynchronization program based on ovarian structures present at the time of NPD (hereafter referred to as short Resynch) would reduce time to pregnancy when compared with blanket use of the d 32 Resynch protocol. Time to pregnancy would be reduced because of the shorter interbreeding interval for CL cows and increased P/AI for no-CL cows. Therefore, the objective of this experiment was to evaluate the effect of short Resynch on time to pregnancy after the first service, P/AI, and physiological outcomes before TAI.

**MATERIALS AND METHODS**

This experiment was conducted from February 2016 to May 2017 on 2 commercial dairy farms located in Tompkins and Cayuga counties in New York. All procedures were approved by the Animal Care and Use Committee of the College of Agriculture and Life Sciences at Cornell University (Ithaca, NY).

On both farms, cows were housed in freestall barns and were fed a TMR once a day with ad libitum access to feed and water. Farm A milked approximately 1,300 cows, with an average milk yield of approximately 43 kg/d. Cows were milked 3 times per day at approximately 8-h intervals until February 2017, when milking frequency changed to 4 times per day at approximately 6-h intervals. All cows received recombinant bST (500 mg of sometribave zinc; Posilac, Elanco Animal Health, Indianapolis, IN) at 10- or 11-d intervals beginning at 80 ± 3 DIM in primiparous cows and 110 ± 3 DIM in multiparous cows until dry-off. Primiparous cows received first service at 82 ± 3 DIM and multiparous cows received first service at 67 ± 3 DIM after synchronization of ovulation with the Double Ovsynch protocol (Souza et al., 2008). Farm B milked approximately 1,900 cows 3 times per day at approximately 8-h intervals and had an average milk yield per cow of approximately 42 kg/d. All cows received recombinant bST at 14-d intervals beginning at 60 ± 3 DIM until dry-off. Primiparous and multiparous cows were synchronized with the Presynch-Ovsynch protocol (Moreira et al., 2001). Cows were eligible to receive AI after the first and second PGF$_{2\alpha}$ treatments of Presynch-Ovsynch at 53 ± 3 and 67 ± 3 DIM, respectively, whereas cows not detected in estrus received TAI at 79 ± 3 DIM. During the experiment, 86.5% (1,103/1,274) of the cows enrolled received the first service at detected estrus, whereas the remaining 13.4% (171/1,274) of the cows received TAI.

Every week, cows that received a previous AI service were blocked by parity (primiparous vs. multiparous) and randomly assigned to the d 32 Resynch (R32) or...
short Resynch (SR) treatments. Cows remained in the same treatment until the end of the experiment. Cows that received their first service postpartum at or after the beginning of the experiment (R32 = 1,010; SR = 1,000) were included for the evaluation of time to pregnancy. Conversely, cows that had already received their first service before the beginning of the experiment were enrolled, but only data from individual TAI services were collected (R32 = 207; SR = 270) for subsequent analysis of P/AI and pregnancy loss. All cows detected in estrus after a previous AI were immediately inseminated. On farm A, detection of estrus was conducted using a combination of visual observation and physical activity monitoring with leg-mounted activity tags (Afi-ActII, Afikim, Kibbutz Afikim, Israel), whereas on farm B, detection of estrus was conducted through visual observation and tail paint removal. Nonpregnancy diagnosis was performed by transrectal ultrasonography (TUS; Ibex Pro, Ibex, Loveland, CO) 32 ± 3 d after AI in cows from both experimental treatments not previously inseminated at detected estrus. In nonpregnant cows, size of follicles and corpora lutea present at NPD was estimated using the ultrasound machine goggle’s screen grid lines comprising squares of 10 mm by 10 mm. Pregnant cows received no further treatment. All TUS examinations were conducted by veterinarians. On farm A the veterinarian was one of the coauthors, whereas on farm B the veterinarians were one of the coauthors and the practicing veterinarian.

Nonpregnant cows from the SR experimental treatment were classified and then treated based on the ovarian structures present at NPD (Figure 1). Cows with at least 1 CL ≥15 mm and at least 1 ovarian follicle ≥10 mm in diameter (CL cows) received PGF$_{2α}$ (500 µg of cloprostenol sodium; Estrumate, Merck Animal Health, Summit, NJ) immediately after NPD, whereas on farm B, detection of estrus was conducted through visual observation and tail paint removal. Nonpregnancy diagnosis was performed by transrectal ultrasonography (TUS; Ibex Pro, Ibex, Loveland, CO) 32 ± 3 d after AI in cows from both experimental treatments not previously inseminated at detected estrus. In nonpregnant cows, size of follicles and corpora lutea present at NPD was estimated using the ultrasound machine goggle’s screen grid lines comprising squares of 10 mm by 10 mm. Pregnant cows received no further treatment. All TUS examinations were conducted by veterinarians. On farm A the veterinarian was one of the coauthors, whereas on farm B the veterinarians were one of the coauthors and the practicing veterinarian.

Nonpregnant cows in the R32 treatment were immediately enrolled in the d 32 Resynch protocol (GnRH, PGF$_{2α}$, 7 d later, GnRH 56 h later, and TAI 16 to 18 h later; Figure 1) to receive TAI. A combination of AI at detected estrus and TAI after the d 32 Resynch protocol without differential treatment based on ovarian structures was selected as a control treatment because it is one of the most commonly used strategies to manage second and greater AI services in lactating dairy cows in the United States (Caraviello et al., 2006; Ferguson and Skidmore, 2013; Scott, 2016).

Reconfirmation of pregnancy was performed 63 ± 3 d after AI by TUS in both farms. A cow was considered to have undergone pregnancy loss when confirmed pregnant at the initial examination and nonpregnant at the time of reconfirmation. Cows with an insemination after a detected estrus between the 2 pregnancy examinations were also considered to have suffered pregnancy loss. Cows diagnosed pregnant 32 ± 3 d after AI but not pregnant at the time of pregnancy reconfirmation (i.e., 63 ± 3 d after AI) were immediately enrolled in the Ovsynch protocol. Cows with detected pregnancy loss failing to conceive after TAI with the Ovsynch protocol were resynchronized after NPD per their experimental treatment (i.e., R32 or SR).

**TUS and Blood Sample Collection for Monitoring Ovarian Responses**

Ovarian responses were monitored at the time of NPD and the GnRH treatment before TAI in a subgroup of cows from each treatment (153 and 146 in R32 and SR, respectively) on farm A. Transrectal ultrasonography was conducted to record size (diameter) of the largest follicle and corpora lutea present on the ovaries. Size of follicles and corpora lutea was estimated using the ultrasound machine (same used for NPD) goggle screen grid lines comprising squares of 10 mm by 10 mm. If the largest follicle was greater than 25 mm and thereby a potential follicular cyst, the second largest follicle was considered the dominant follicle for analysis. From the same group of cows, blood samples were collected to determine circulating concentration of P4. Blood samples were collected using 8-mL heparinized evacuated tubes (Vacutainer, Becton Dickinson, Franklin Lakes, NJ) via puncture of the coccygeal vein or artery. Samples were placed in a cooler with ice until being transported to the laboratory within ≤4 h of collection. Samples were centrifuged at 1,700 × g for 20 min at 4°C. Plasma aliquots were harvested and transferred to vials for storage at −20°C until assays were performed. Progesterone concentration data at NPD were available for 205 cows with a CL ≥15 mm and 78 cows with no
CL ≥15 mm at NPD from both treatments. Data from ovarian structures and P4 concentration at the GnRH treatment before TAI was available for 114 and 121 cows from the R32 and SR treatments, respectively. The proportion of cows with a functional CL based on circulating concentration of P4 (P4 ≥1 ng/mL) was the outcome of interest at NPD. At the time of the GnRH injection before TAI, the outcomes of interest were (1) the proportion of cows with low P4 based on a cutoff value of ≤0.5 ng/mL and (2) the size of the largest ovarian follicle present. The cutoff for circulating P4 concentration at the time of the GnRH treatment before TAI was selected based on recent data suggesting that P4 concentrations below 0.5 to 0.4 ng/mL at this time point are associated with greater P/AI than P4 levels above such cutoff points (Brusveen et al., 2009; Giordano et al., 2012c).

**Determination of P4 Concentration**

Concentration of P4 in plasma was determined in duplicate with a commercial solid-phase, no-extraction RIA (ImmuChem coated tube, MP Biomedicals, Costa Mesa, CA). To assess precision of the assays, control samples with high (5.8 ng/mL) and low (0.3 ng/mL) concentrations of P4 were included at the beginning and end of each assay (n = 6 assays). Average detection limit for the P4 assay was 0.1 ng/mL. Average intra-assay coefficient of variation (CV) for the high-concentration sample was 9.7%, whereas the interassay CV was 13.4%. For the low-concentration sample the average intra-assay CV was 24.2%, whereas the interassay CV was 31.3%.

**Statistical Analysis**

This experiment was conducted as a complete randomized block design using parity (primiparous vs. multiparous) as blocking factor. According to sample size calculations conducted using the sample size calculation option of WinPepi version 11.51 (Abramson, 2011), a total of 398 cows per treatment was needed to detect a hazard ratio for pregnancy of 1.25 with an average probability of survival at the end of the experimental period of 20%, probability of type I error rate of 5%, and probability of type II error rate of 20%.
Cox’s proportional hazards analysis for pregnancy after the first service postpartum was conducted using the PHREG procedure of SAS (version 9.4, SAS Institute Inc., Cary, NC), with treatment, parity, and milk production tercile (calculated for farm and parity group within farm) as fixed effects and farm as random effect. Milk production level was removed from the final model because $P > 0.10$. Only cows that did not conceive to first AI and that had at least 210 d at risk of becoming pregnant after the first service were included in the analysis. A cow was considered pregnant for the analysis of time to pregnancy only if it was reconfirmed pregnant 63 ± 3 d after AI. Cows that became not eligible for AI due to a farm management decision (i.e., coded as “do not breed”) or that left the herd due to sale or death were right censored. Kaplan–Meier survival curves were generated to illustrate the rate of pregnancy after the first service postpartum using the survival analysis option of MedCalc (version 12.5.0.0; MedCalc Software, Ostend, Belgium).

Binary outcomes (i.e., $\text{P/AI at 32 ± 3 and 63 ± 3 d after AI}$, pregnancy loss, proportion of cows inseminated at detected estrus, proportion of cows not pregnant 210 d after the first service, and proportion of cows with high or low P4 concentration at NPD and the GnRH before TAI) were analyzed using logistic regression with the GLIMMIX procedure of SAS. Pregnancy per AI at 32 ± 3 d after AI was evaluated for AI services at detected estrus, TAI services, and both types of AI services combined, whereas $\text{P/AI at 63 ± 3 d after AI and pregnancy loss were evaluated for AI services at detected estrus and TAI combined.}$

Treatment and parity (primiparous vs. multiparous) were included as fixed effects and farm as random effect. Days in milk at insemination ($\leq 100$, $>100$ and $\leq 150$, $>150$ and $\leq 200$, and $>200$ d) and season of insemination [warm (June to August) vs. cold (September to May)] were offered to the initial models for $\text{P/AI and pregnancy loss.}$ Season was removed from all final models and DIM was removed from the pregnancy loss models because $P > 0.10$.

The level of agreement between TUS and P4 concentration (reference test) to detect the presence of an active CL was determined through calculation of the kappa value for interrater agreement obtained with the FREQ procedure of SAS. Size of the largest ovarian follicle at the time of GnRH before TAI was analyzed by ANOVA using the MIXED procedure of SAS, including treatment and parity as fixed effects.

All proportions reported were generated using the FREQ procedure of SAS, whereas values for quantitative outcomes are reported as arithmetic means calculated using the MEANS procedure of SAS. All explanatory variables were considered significant if $P \leq 0.05$, whereas $P > 0.05$ and $\leq 0.10$ was considered a tendency.

**RESULTS**

**Time to Pregnancy After First Service**

In total, 485 and 462 cows from the R32 and SR treatments, respectively, were not pregnant after the first service postpartum and were at least 210 d at risk of becoming pregnant after the first service postpartum unless they left the herd due to sale or death. The proportion of second and greater AI services that occurred after estrus detection [$P = 0.56; \text{R32} = 66.4\% (739/1,113) \text{vs. SR} = 67.7\% (594/878)$] and P/AI for these services [$P = 0.26; \text{R32} = 34.4\% (277/805) \text{vs. SR} = 37.2\% (245/658)$] was similar for both treatments.

The hazard of pregnancy was greater ($P = 0.03$) for the SR treatment than for the R32 treatment (hazard ratio = 1.18; 95% CI: 1.01–1.37; Figure 2) and was similar ($P = 0.15$) for primiparous and multiparous cows (hazard ratio = 1.12; 95% CI: 0.96–1.31). Median time to pregnancy was 95 d (95% CI: 84–108) and 79 d (95% CI: 71–96) for the R32 and SR treatments, respectively. Mean time to pregnancy was 111 ± 3 and 100 ± 3 d for the R32 and SR treatments, respectively. At 210 d after first service, a greater ($P < 0.02$) proportion of cows were not pregnant in the R32 treatment (29.3%; 142/485) than in the SR treatment (22.3%; 103/462).

![Figure 2](https://example.com/figure2.png)
and a greater \((P < 0.01)\) proportion of multiparous cows (29.3%; 177/605) than primiparous cows (19.9%; 68/342) were not pregnant.

**P/AI and Pregnancy Loss**

At NPD the proportion of CL cows was 71.3\% (513/720) for the R32 treatment and 71.2\% (583/819) for the SR treatment. The rest of the cows were classified as no-CL cows in both treatments. The proportion of cows inseminated at detected estrus after NPD and before TAI was greater for the R32 treatment for all cows \([P < 0.001]; \text{R32} = 14.6\% (105/720); \text{SR} = 8.7\% (71/819)\) and cows with a CL at NPD \([P < 0.001]; \text{R32} = 14.4\% (74/513); \text{SR} = 6.5\% (38/583)\). No effect of treatment \(P = 0.68\) was observed for cows with no CL at NPD \([\text{R32} = 15.0\% (31/207); \text{SR} = 14.0\% (33/236)]\). For the R32 treatment, 50.9\% of the AI services at detected estrus were conducted from the time of the first GnRH until the PGF2α injection of the protocol, whereas the remaining 49.1\% were conducted after the PGF2α and before the morning of TAI.

Overall, P/AI at 63 ± 3 d after TAI for all AI services (CL and no-CL cows combined) was not affected by treatment \([P = 0.50]; \text{R32} = 27.8\% (198/713); \text{SR} = 29.1\% (233/801)\) or parity \([P = 0.27]; \text{primiparous} = 29.6\% (184/621); \text{multiparous} = 27.7\% (247/893)\). Among cows with a CL at NPD there was no effect of treatment \([P = 0.40]; \text{R32} = 29.9\% (152/508); \text{SR} = 27.4\% (156/569)\) or parity \([P = 0.12]; \text{primiparous} = 31.0\% (135/436); \text{multiparous} = 27.0\% (173/641)\) on P/AI. For cows with no CL at NPD, P/AI was greater \((P = 0.01)\) for the SR treatment \((33.2\%; 77/232)\) than for the R32 treatment \((22.4\%; 46/205)\), but there was no effect of parity \([P = 0.82]; \text{primiparous} = 26.5\% (49/185); \text{multiparous} = 29.4\% (74/252)\). Days in milk at TAI affected P/AI 63 ± 3 d after AI for all AI services \((P = 0.03)\) and for cows with no CL at NPD \((P < 0.001)\) because cows with DIM >200 had reduced \((P < 0.05)\) P/AI compared with cows inseminated at earlier DIM. For AI services in cows with a CL at NPD, P/AI was not affected \((P = 0.93)\) by treatment for all inseminations, inseminations after a detected estrus \((P = 0.21)\), or TAI services \((P = 0.94); \text{Table 1}\). For AI services in cows with no CL at NPD, P/AI was greater for the SR treatment than for the R32 treatment for all inseminations \((P = 0.02)\) and TAI services \((P < 0.01)\), but it did not differ for inseminations after a detected estrus \((P = 0.58); \text{Table 1}\).

Overall, P/AI at 63 ± 3 d after TAI for all AI services (CL and no-CL cows combined) was not affected by treatment \([P = 0.50]; \text{R32} = 27.8\% (198/713); \text{SR} = 29.1\% (233/801)\) or parity \([P = 0.27]; \text{primiparous} = 29.6\% (184/621); \text{multiparous} = 27.7\% (247/893)\). Among cows with a CL at NPD there was no effect of treatment \([P = 0.40]; \text{R32} = 29.9\% (152/508); \text{SR} = 27.4\% (156/569)\) or parity \([P = 0.12]; \text{primiparous} = 31.0\% (135/436); \text{multiparous} = 27.0\% (173/641)\) on P/AI. For cows with no CL at NPD, P/AI was greater \((P = 0.01)\) for the SR treatment \((33.2\%; 77/232)\) than for the R32 treatment \((22.4\%; 46/205)\), but there was no effect of parity \([P = 0.82]; \text{primiparous} = 26.5\% (49/185); \text{multiparous} = 29.4\% (74/252)\). Days in milk at TAI affected P/AI 63 ± 3 d after AI for all AI services \((P = 0.03)\) and for cows with no CL at NPD \((P < 0.001)\) because cows with DIM >200 had reduced \((P < 0.05)\) P/AI compared with cows inseminated at earlier DIM. On the other hand, DIM did not affect \((P = 0.52)\) P/AI for cows with a CL at NPD.

Pregnancy loss for all AI services combined (CL and no-CL cows combined) was similar for both treatments \([P = 0.32]; \text{R32} = 8.3\% (18/216); \text{SR} = 10.4 (27/260)\) and was not affected by parity \([P = 0.68]; \text{primiparous} = 29.3\% (177/605); \text{primiparous} = 19.9\% (68/342)\) were not pregnant.

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**Table 1. Pregnancy per AI and pregnancy loss for cows that received the experimental treatments**

<table>
<thead>
<tr>
<th>Item</th>
<th>R32 [% (no./no.)]</th>
<th>SR [% (no./no.)]</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All inseminations for CL and no-CL cows combined</td>
<td>31.0 (223/720)</td>
<td>33.9 (278/819)</td>
<td>0.17</td>
</tr>
<tr>
<td>AI at detected estrus</td>
<td>39.1 (41/105)</td>
<td>43.7 (31/71)</td>
<td>0.57</td>
</tr>
<tr>
<td>TAI</td>
<td>29.6 (182/615)</td>
<td>33.0 (247/748)</td>
<td>0.14</td>
</tr>
<tr>
<td>All inseminations for CL cows</td>
<td>32.7 (168/513)</td>
<td>32.8 (191/583)</td>
<td>0.93</td>
</tr>
<tr>
<td>AI at detected estrus</td>
<td>37.8 (28/74)</td>
<td>50.0 (19/38)</td>
<td>0.21</td>
</tr>
<tr>
<td>TAI</td>
<td>31.9 (140/439)</td>
<td>31.6 (172/545)</td>
<td>0.94</td>
</tr>
<tr>
<td>All inseminations for no-CL cows</td>
<td>28.6 (55/207)</td>
<td>36.9 (87/236)</td>
<td>0.02</td>
</tr>
<tr>
<td>AI at detected estrus</td>
<td>41.9 (13/31)</td>
<td>36.4 (12/33)</td>
<td>0.58</td>
</tr>
<tr>
<td>TAI</td>
<td>23.9 (42/176)</td>
<td>37.0 (75/203)</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

1Cows were randomly assigned to the d 32 Resynch (R32) or short Resynch (SR) treatments. Nonpregnancy diagnosis was performed 32 ± 3 d after AI. Nonpregnant cows in the R32 treatment received the Ovsynch protocol [GnRH, PGF2α, 7 d later, GnRH 56 h later, timed AI (TAI) 16 to 18 h later] for resynchronization of ovulation, whereas cows in the SR treatment were resynchronized based on the ovarian structures present at nonpregnancy diagnosis. Cows with at least 1 corpus luteum ≥15 mm and 1 follicle ≥10 mm received PGF2α, PGF2α 24 h later, GnRH 32 h later, and TAI 16 to 18 h after GnRH. Cows that did not fulfill the corpus luteum and follicle size criteria received a modified Ovsynch protocol with 2 PGF2α treatments and progesterone supplementation (GnRH plus CIDR, PGF2α, and CIDR removal 7 d later, PGF2α 24 h later, GnRH 32 h later, and TAI 16 to 18 h later).

2Cows with at least 1 corpus luteum ≥15 mm and 1 follicle ≥10 mm.

3Cows with no corpus luteum ≥15 mm or no follicle ≥10 mm.

DIM = Days in Milk; TAI = Time of Artificial Insemination; CL = Corpus Luteum; PGF2α = Prostaglandin F2α; GnRH = Gonadotropin Releasing Hormone; CIDR = Continuous Intramammary Device Replacement; P/AI = Pregnancy per Artificial Insemination; NPD = Nonpregnancy Diagnosis; AI = Artificial Insemination.
A SHORT RESYNCH PROGRAM IMPROVES PREGNANCY RATE

= 8.9% (18/202); multiparous = 9.6% (27/274)]. For cows with a CL at NPD, pregnancy loss tended (P = 0.06) to be greater for the SR treatment (11.9%; 21/177) than for the R32 treatment (6.8%; 11/163), but there was no effect of parity [P = 0.87; primiparous = 8.9% (13/148); multiparous = 9.9% (19/192)]. For cows with no CL at NPD, treatment (P = 0.25) and parity (P = 0.84) did not affect pregnancy loss [R32 = 13.2% (7/53) vs. SR = 7.2% (6/83); primiparous = 9.3% (5/54) vs. multiparous = 9.8% (8/82)].

Ovarian Structures and P4 Concentration at NPD

Among cows with a CL ≥15 mm, 84.4% had P4 ≥1 ng/mL, 3.9% had P4 >0.5 and <1 ng/mL, and 11.7% had P4 ≤0.5 ng/mL. Among cows with no CL ≥15 mm at NPD, 29.5% had P4 ≥1 ng/mL, 51.1% had P4 >0.5 and <1 ng/mL, and 19.4% had P4 ≤0.5 ng/mL. Among cows with no CL ≥15 mm and P4 ≥1 ng/mL at NPD, 43.5% (10/23) had at least 1 fluid-filled cavity ≥25 mm (usually considered an ovarian cyst) present on their ovaries. The agreement between TUS and P4 concentration to determine the presence of a functional CL (P4 ≥1 ng/mL) was moderate (kappa = 0.53; 95% CI: 0.42–0.64; P < 0.001).

Ovarian Structures and P4 Concentration at the GnRH Treatment Before TAI

Overall, a greater (P = 0.02) proportion of cows had P4 concentration <0.5 ng/mL in the SR treatment (81.8%; 99/121) than in the R32 treatment (68.4%; 78/114). In addition, for cows with a CL at NPD, the proportion of cows with P4 ≤0.5 ng/mL tended (P = 0.08) to be greater for the SR treatment than for the R32 treatment (Table 2). There was no effect of treatment (P = 0.14) for cows without a CL at NPD (Table 2).

Cows in the R32 treatment tended (P = 0.10) to have larger follicles (18.0 ± 0.4 mm) than cows in the SR treatment (17.1 ± 0.4 mm). For cows with a CL at NPD, treatment did not affect (P = 0.74) the size of the largest follicle at the time of GnRH (Table 2), whereas for cows with no CL at NPD, follicle size at GnRH treatment tended (P = 0.07) to be greater for cows in the R32 treatment (Table 2).

DISCUSSION

A program (short Resynch) that relied on resynchronization of ovulation treatments given to cows based on the ovarian structures present at NPD was compared with blanket use of the d 32 Resynch protocol. In support of our hypothesis, cows in the SR program had fewer days to pregnancy after the first service. Another benefit of the SR treatment was a reduction in the proportion of nonpregnant cows at the end of the experimental period. Thus, the SR treatment may benefit dairy herds when compared with blanket use of the d 32 Resynch protocol through a reduction in both time to pregnancy and the proportion of nonpregnant cows removed from the herd. Of note, removal of the GnRH treatment 25 ± 3 d after AI eliminated 2 unintended consequences: (1) interfering with estrus expression before NPD and (2) unnecessary treatment of pregnant cows with GnRH. In addition, we observed that cows with a CL ≥15 mm and a follicle ≥10 mm at NPD that received the short resynchronization protocol had a P/TAI similar to that of cows with the same ovarian status that received the d 32 Resynch protocol. Conversely, no-CL cows clearly benefited from P4 supplementation and the 2 PGF2α treatments.

### Table 2. Ovarian status at the time of GnRH treatment before timed AI for cows that received the experimental treatments

<table>
<thead>
<tr>
<th>Item</th>
<th>CL ≥15 mm and follicle ≥10 mm</th>
<th>No CL ≥15 mm or no follicle ≥10 mm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R32 (n = 80)</td>
<td>SR (n = 88)</td>
</tr>
<tr>
<td>Progesterone ≤0.5 ng/mL [% (no.)]</td>
<td>70.0 (56)</td>
<td>81.8 (72)</td>
</tr>
<tr>
<td>Size of the largest follicle (mm)</td>
<td>17.1 ± 0.5</td>
<td>16.8 ± 0.4</td>
</tr>
</tbody>
</table>

1Cows were randomly assigned to the d 32 Resynch (R32) or short Resynch (SR) treatments. Nonpregnancy diagnosis was performed 32 ± 3 d after AI. Nonpregnant cows in the R32 treatment received the Ovsynch protocol [GnRH, PGF2α, 7 d later, GnRH 56 h later, timed AI (TAI) 16 to 18 h later] for resynchronization of ovulation, whereas cows in the SR treatment were resynchronized based on the ovarian structures present at nonpregnancy diagnosis. Cows with at least 1 corpus luteum ≥15 mm and 1 follicle ≥10 mm received PGF2α, PGF2α 24 h later, GnRH 32 h later, and TAI 16 to 18 h after GnRH. Cows that did not fulfill the corpus luteum and follicle size criteria received a modified Ovsynch protocol with 2 PGF2α treatments and progesterone supplementation (GnRH plus CIDR, PGF2α, and CIDR removal 7 d later, PGF2α 24 h later, GnRH 32 h later, and TAI 16 to 18 h later).

2NPD = nonpregnancy diagnosis.

3CL = corpus luteum.
Various strategies for resynchronization of ovulation to maximize P/AI (Dewey et al., 2010; Giordano et al., 2012b, c), shorten the interval between TAI services (Fricke et al., 2003; Galvão et al., 2007; Sinedino et al., 2014), or maximize the number of cows inseminated upon estrus detection (Bruno et al., 2013; Chebel et al., 2013; Giordano et al., 2015) have been developed and tested. Nonetheless, few experiments evaluated the effect of these strategies on time to pregnancy, which is the ultimate driver of dairy cow profitability. This is important when the interbreeding interval or the resynchronization dynamics (i.e., proportion of cows undergoing AI at detected estrus vs. TAI) are different for the groups compared. Indeed, cows in the SR treatment had reduced time to pregnancy, primarily due to the 7-d shorter interbreeding interval for the approximately 70% of cows with a CL at NPD despite no gain in P/AI. The greater P/AI without extending the interbreeding interval for no-CL cows in the SR treatment also contributed to reduced time to pregnancy; however, only approximately 30% of the total nonpregnant cows benefited from greater P/AI.

The similar P/AI for CL cows in both treatments was critical for reducing time to pregnancy for SR because if cows that received this protocol had reduced P/AI, the benefit of shorter interbreeding interval could have been counterbalanced by the reduced proportion of pregnant cows after TAI. For CL cows in the SR treatment, the detriment of suboptimal control of the follicular wave dynamics (Wijma et al., 2017) must have been compensated, at least in part, by the 2 PGF2α treatments. Indeed, the proportion of cows with a CL at NPD that had low P4 concentration at the time of GnRH treatment before TAI tended to be greater in the SR treatment. We speculate that maximizing the proportion of cows with low P4 concentration at the time of inducing ovulation before TAI had a greater effect on P/AI than reduced control of the follicular wave from which the ovulatory follicle emerged. In this regard, previous experiments with resynchronized lactating dairy cows reported greater P/AI for cows that failed to ovulate to the first GnRH treatment of Ovsynch but had complete luteolysis before TAI than for cows that ovulated but had incomplete luteolysis before TAI (Giordano et al., 2012c). Other potential issues with synchronization of ovulation protocols that do not tightly control follicular wave dynamics before TAI include lack of ovulation to the GnRH before TAI and the potential to increase pregnancy losses due to poor oocyte quality or luteal insufficiency after AI. Cows in which the follicular wave of the ovulatory follicle began less than 5 d before induction of luteolysis may have either failed to ovulate in response to the GnRH before TAI (Vasconcelos et al., 1999) or ovulated a small follicle that resulted in a small CL incapable of supporting pregnancy (Vasconcelos et al., 2001). Cows that initiated the follicular wave of the ovulatory follicle more than 5 d before induction of luteolysis may have been more likely to have poor embryo quality because of ovulation of an aged oocyte (Cerri et al., 2009). These issues were expected for some CL cows in the SR treatment because GnRH was not given to induce a follicular wave from which the ovulatory follicle would emerge. In this regard, we recently reported greater variation for age of the largest follicle and greater follicle size at NPD 32 ± 3 d after AI in cows that did not receive GnRH compared with cows that received GnRH 7 d earlier (Wijma et al., 2017).

We speculate that the tendency for greater pregnancy loss in the SR treatment for CL cows could be explained, at least in part, by reduced control of the follicular wave dynamics before TAI. In our experiment, however, differences in pregnancy loss between groups may have also been less dramatic because the follicular wave dynamics of cows resynchronized with Ovsynch-type protocols is not optimal due to ovulatory failure in response to the first GnRH of the protocol in a high proportion of previously inseminated cows (Giordano et al., 2012c; Wijma et al., 2017). In particular, poor ovulatory response has been reported for cows with a functional CL at the time of the GnRH treatment (Galvão and Santos, 2010; Giordano et al., 2012c).

Despite no reduction of the interbreeding interval for no-CL cows, the increment in P/AI for these cows likely contributed to the reduction of time to pregnancy for the SR treatment. Greater P/AI for resynchronized services at the same interbreeding interval reduces time to pregnancy by decreasing the need for resireminations at a later time. In cows with no CL at NPD, the beneficial effect of P4 supplementation and 2 PGF2α treatments likely synergized. Supplemental P4 in cows without a CL at the initiation of Ovsynch-type protocols has been proven effective for increasing P/AI (Bartolome et al., 2009; Bisinotto et al., 2013, 2015) due to a combination of (1) improved embryo quality because of improved endocrine environment for oocyte development (Rivera et al., 2011), (2) improved uterine environment (Cerri et al., 2011), and (3) reduced premature estrus and ovulation before TAI (Stevenson et al., 2006). The latter, however, may not have been as relevant in the current experiment because cows detected in estrus any time were immediately inseminated and had reasonable P/AI. An additional PGF2α treatment after induction of luteolysis is known to increase the proportion of cows with complete luteolysis before TAI (Brusveen et al., 2009; Ribeiro et al., 2012; Wiltbank et al., 2015), which in turn increases P/AI (Souza et al., 2007; Brusveen et al., 2009). In our experiment, the additional PGF2α treatment may explain a substan-
tial portion of the gain in P/AI for no-CL cows. These cows are more likely to ovulate after GnRH (Galvão and Santos, 2010; Giordano et al., 2012c; Lopes et al., 2013) due to a greater GnRH-induced LH response (Giordano et al., 2012a; Lima et al., 2013; Pulley et al., 2015) and thereby are more likely to have only one 6-d-old CL not fully responsive to PGF2α at the time of induction of luteolysis. That may explain the relatively low proportion of cows with low P4 at the time of the GnRH before TAI even after 2 PGF2α treatments. Our low proportion of cows with low P4 at the time of the induction of luteolysis would have allowed the CL to receive the treatment for no-CL cows. In particular, the subgroups of cows likely benefited from receiving the modified Ovsynch protocol with P4 supplementation.

Although our control group could have easily incorporated the modified protocol with 2 PGF2α treatments for all cows and P4 supplementation for no-CL cows, our reason for not doing so was to keep the treatment as simple as possible and reflect the conditions of many dairy herds in the United States that use this strategy for resynchronization of ovulation (Caraviello et al., 2006; Ferguson and Skidmore, 2013; Scott, 2016). Most farms likely choose to either not treat cows based on their ovarian status at NPD 32 ± 3 d after AI or conduct NPD 39 ± 3 d coincident with the time of the PGF2α treatment before TAI. Whether differential treatment of no-CL cows in the R32 treatment would have offset the gain in reproductive performance from the 7-d shorter interbreeding interval in the SR treatment is unknown. Nevertheless, it is rather unlikely because most of the cows had a CL at NPD; therefore, they benefited by the reduced interbreeding interval of the SR protocol.

Finally, another important consideration for the use of the SR treatment as a management strategy is the potential extra cost of the program compared with blanket use of d 32 Resynch. The expected reduction in time to pregnancy and proportion of cows not pregnant in late lactation must offset the additional cost of the protocol for no-CL cows and the potential extra cost of TUS in farms that do not routinely use this method of NPD. Thus, the economics of more complex and potentially more expensive management strategies such as SR warrant further investigation.

CONCLUSIONS

A reproductive management program designed to (1) reduce the interbreeding interval for TAI services in cows with a CL ≥15 mm and a follicle ≥10 mm at NPD and (2) increase P/AI of cows without a CL at NPD through a modified Ovsynch protocol with 2 PGF2α treatments and P4 supplementation reduced time to pregnancy and the proportion of nonpregnant cows 210 d after first service compared with blanket use of the d 32 Resynch protocol. Therefore, the SR program is a new management strategy with the potential to improve reproductive performance of dairy herds that enroll nonpregnant cows not reimplanted at detected estrus in Ovsynch-type resynchronization of ovulation protocols regardless of their ovarian status. In addition,
removing the GnRH treatment to induce a new follicular wave in the SR treatment helped reduce the interbreeding interval without disrupting estrus behavior and reduced unnecessary treatment of pregnant cows with GnRH 25 ± 3 d after AI.

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REFERENCES


