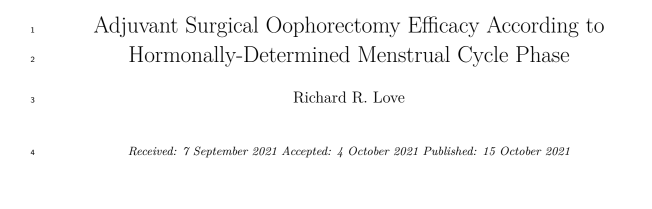


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6 Abstract

Purpose: While there is now considered to be no significant outcome impact of the timing of 7 breast surgery in the menstrual cycle of premenopausal women with breast cancer, the data with respect to adjuvant surgical oophorectomy in women with breast cancer have received 9 limited exposition and attention. In a trial investigating the timing of surgical ophorectomy 10 in women with metastatic disease, we observed a trend for poorer overall survival in women in 11 women in prolonged follicular phases of the menstrual cycle, with low progesterone levels. 12 Methods: The data from a previously reported adjuvant randomized clinical trial addressing 13 the timing of surgical ophorectomy in the menstrual cycle have been examined in detail, 14 presenting here new data from pre-planned secondary analyses. Multivariable Cox models 15 were used. Methods: The data from a previously reported adjuvant randomized clinical trial 16 addressing the timing of surgical ophorectomy in the menstrual cycle have been examined in 17 detail, presenting here new data from pre-planned secondary analyses. Multivariable Cox 18

¹⁹ models were used.

21 Index terms— adjuvant therapy, surgical oophorectomy, tamoxifen, menstrual cycle timing.

22 **1 I**.

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Background lobally, 500,000 premenopausal women annually present with hormone receptor positive breast 23 24 cancer. For these women with operable disease, surgical oophorectomy or ovarian function-suppression plus 25 tamoxifen are the most effective adjuvant therapies [1,2,3]. Secondary analysis of women in a clinical trial receiving surgical ophorectomy treatment suggested that if the ophorectomy surgery was performed during 26 the luteal phase of the menstrual cycle, long term disease-free and overall survival were significantly better than 27 if the surgery was done in the follicular phase [4]. We have conducted and reported two phase III trials, one 28 in metastatic and one in adjuvant patients, to investigate this finding in which we presented some data from 29 secondary pre-planned analyses of outcomes according to hormonally confirmed menstrual cycle phases [5,6]. 30

In the reported metastatic study, the primary analysis showed that the randomized luteal history (beyond day 14 since beginning of last menstrual period) and follicular history (from beginning day of menstrual period through day 14) surgical oophorectomy patients had equivalent overall survival (L H =F H for OS) [6]. In pre-planned analyses of all randomized patients with hormonal levels, based on confirmed hormonal status L H patients with high progesterone (Pg) levels had better overall survival than L H patients with low progesterone levels: 27 versus 17 months (multivariable p=0.14) [6].

The primary analysis of the adjuvant trial showed that luteal phase by history(L H)patients, did not This communication reports new data from the adjuvant study, other data, and interpretations relevant to our findings. have better survival than patients in historical follicular phase, F H, by strong trends (multivariable overall survival p=0.05) [5, ??igure 2]. That is, contrary to the study hypothesis, L H patients had worse disease-free (DFS) and overall survival. One exploratory analysis result was presented: In patients randomized to receive mid-luteal phase surgery, patients with higher Pg (?2ng/ml) had better DFS than those with < 2ng/ml (aHR 0.53; 95% CI 0.34 -0.84; p=0.006) [5].

44 **2** II.

$_{45}$ 3 Methods

Reports of two phase III clinical trials of surgical oophorectomy plus tamoxifen (SO +T) in adjuvant and 46 metastatic populations have been published with the detailed designs, eligibilities, IRB approvals, treatments, 47 laboratory studies and statistical methods [5,6]. A G consort figure for the adjuvant trial populations that are 48 the subject of this report is presented in figure 1. In this study, 383 patients (of 509 randomized because they 49 would not be by history in luteal phase-that is beyond day 14 since last menstrual period began-for the next 1-6 50 days) had: 1. menstrual cycle history data; 2. day-ofsurgery blood hormone level determinations showing levels 51 of <2ng/ml or 5 or greater ng/ml; and 3. complete follow up data. In the current report, Cox model subgroup 52 analyses are based on data from these 383 subjects. In this adjuvant trial are three subgroups of the combined two 53 randomized groups, defined by menstrual cycle dates history and hormonal levels on the dates of oophorectomy 54 surgery (Figure 1). In this report luteal phase history patients with progesterone levels of >=2 but < 5 ng/ml have 55 been removed to provide information on the most well-defined follicular and luteal groups. The three subgroups 56 of patients are: Follicular phase patients-by-history with progesterone levels <2ng.ml-"F H confirmed"; luteal 57 58 phase-by-history patients with progesterone levels = >5 ng/ml-"L H confirmed"; and luteal phase-by-history 59 patients with progesterone levels <2 ng/ml or prolonged follicular phase patients, or anovulatory patients-"L H 60 unconfirmed". If the less well-defined subgroup of 49 luteal phase by history patients with progesterone levels of between 2 and 5 ng/ml, half of whom were in follicular phase by history, is included as confirmed luteal phase 61 62 patients, the results reported here are unchanged.

A multivariable Cox proportional hazards model was used to estimate adjusted hazard ratios between pairs of luteal phase-confirmed and unconfirmed, and luteal phase-unconfirmed and follicular phaseconfirmed groups. In these analyses, the other prognostic variables included were: adjuvant radiotherapy, stage, nodal status, tumor size and patient age. As in the report of the primary analyses "proportionality assumptions for the Cox models were assessed by diagnostic plots of the scaled Schoenfeld residuals and log-minus-log survival plots. Substantial deviations from proportionality were not observed."

In all comparisons of these randomized patient subgroups, treatment group assigned at random assignment was compared regardless of the treatment received (5). P values are reported for completeness: because these are exploratory/explicatory analyses, they cannot be considered hypothesis testing results.

72 **4 III.**

$_{73}$ 5 Results

In pre-planned analyses based on historyconfirmed hormonal status, the explanation for the definitive primary
analysis result is clear.[The result described above: luteal phase by history patients, did not have better survival
than patients in historical follicular phase, (multivariable overall survival p=0.05)].

The subgroups of unconfirmed and confirmed luteal phase status had markedly different survival experiences. Among all combined randomized patients, L H patients with high progesterone levels ("L H confirmed", n=150) had better survival than L H patients with low progesterone levels ("L H unconfirmed", n=112): the differences at 5 years were for disease free survival, 20%, HR=1.60 (95% C.I.:1.07-2.38), multivariable p=0.02; and for overall survival,15%, HR=1.63 (95% C.I. 1.03-2.56), multivariable p=0.036. The differences between F H confirmed (n=121) and L H unconfirmed (n=112) for both DFS and OS were marginally greater.

Among all randomized L H patients: those with high progesterone had better survival than those with low progesterone (p=0.001).

85 IV.

⁸⁶ 6 Discussion a) Interpretation

The reported new results show that in preplanned exploratory analyses in a second phase IIIadjuvant study, 87 among the randomized patients, those patients found to be in prolonged follicular phase (that is beyond day 88 14 of their menstrual cycle) with low progesterone levels at the times of their ophorectomy surgeries, showed 89 limited evidence of long-term disease-free and overall survival benefits, despite receiving additionally tamoxifen 90 treatment. A conservative interpretation is that these observations define a new hypothesis. The major limitation 91 of the results is that they are secondary study findings, whose statistical significance cannot be reliably estimated. 92 The major strength of the results is that they have been found among randomized patients in two studies (5,6). 93 94 As I have previously written, which critically bears repetition here: "the corollary to this new observation 95 is that were such unconfirmed luteal phase patients (in these and other studies usually one third of patients) 96 identified a priori, and not treated with this surgery at this time, those patients treated in hormonally-confirmed 97 follicular or luteal phases would be expected to have better outcomes that the average outcomes that are seen from this treatment applied to all premenopausal women regardless of hormonal status and menstrual cycle phase. 98 Thus, if in a high-risk group of women with operable breast cancer receiving SO (+T) (without paying any 99 attention to their menstrual cycle history and blood levels of progesterone), 65% have no recurrence in 5 years; 100 if patients have their SO in the first half of their menstrual cycles by history and with confirmation showing 101 low progesterone blood levels, 72% will have no recurrence in 5 years. This increased level of benefit from 102

appropriately timed SO, suggests that timed SO+T is more effective than GnRH + tamoxifen, and equivalently 103 effective or better than GnRH + aromatase inhibitor" [2]. Further discussion is warranted. The adjuvant therapy 104 primary analysis results are definitive that patients in historical luteal phase are extremely unlikely to have 105 better outcomes than patients in historical follicular phase [5]. The data presentation in the primary publication, 106 while reporting the one exploratory analysis finding of better DFS in confirmed luteal versus unconfirmed luteal 107 patients, was conservative in combining all patients in the trial, randomized and nonrandomized. Because for 108 unexplained reasons the nonrandomized patients enjoyed better-than-expected survival, the striking finding in 109 the randomized patients reported here above, was not found. Differences in outcomes in non-randomized versus 110 randomized groups of patients have been repeatedly observed, explained by selection bias, so these findings are 111 not unusual, and are the basis for the current report emphasizing the clear explicatory findings for the primary 112 trial result, and their consistency with the results of the metastatic trial [6,7]. 113

studies are correct and represent 'truth', given this different definition, theoretically the original study might 114 be expected to show the same result. This is because if we make the assumption that day of surgery in the 115 menstrual cycle is always F H + 6, and L H + 6, new L H defined patients will all be beyond day 21 in their 116 cycles and more likely to be in hormonally-confirmable luteal phase (which patients in the new adjuvant and 117 metastatic studies did well), and new F H patients will include true F patients, and prolonged F patients (or 118 119 "L H unconfirmed"), the latter sub-group of whom did badly in the new studies as discussed above [5,6]. Thus, 120 conceivably the original study could in fact, with appropriate definitions of day one of the cycle, give the same L H (very likely confirmable) better result than in a combined group of F H (likely confirmable) and F H (prolonged) 121 (=L H unconfirmed). When re-analyses were done under these new definitions, no DFS and OS differences were 122 seen between the two redefined L H and F H groups. Given the now-likely poor and mixed patient and physician 123 definitions quality of the menstrual cycle history data in this study, this revised result is not surprising [4]. 124

¹²⁵ 7 b) The hypothesis-generating study data and their interpre tation

The previous hypothesis-generating study also deserves comment [4]. The discussed adjuvant study was designed 127 to test the hypothesis that surgery during historical luteal phase (L H) of the menstrual cycle had superior efficacy 128 [5]. This design followed from secondary exploratory analyses of an adjuvant study of surgical oophorectomy plus 129 tamoxifen, which strongly suggested that L H was superior [4]. How can the findings from these 3 studies be 130 131 reconciled [4,5,6]? The hypothesis-generating study categorized patients as being F H or L H based on reported "day one' of their menstrual cycle at the time of their breast and surgical oophorectomy surgeries (done under 132 133 the same anesthesia on the same days) [4]. Without careful discussion of this time point, we assumed that day 134 one of the menstrual cycle according to the Vietnamese women was the day they began their menstrual bleeding. 135 In discussions with Vietnamese, now American immigrant women, who had resided in Vietnam during the same period the study was conducted and who were in the same age range as the study subjects, these women indicated 136 that their definition of day one of their menstrual cycle when they were in Vietnam, was the day they had no 137 further menstrual bleeding. In exploring this possibility with the 3 Vietnamese investigating physicians, they 138 agreed that this misunderstanding was very plausible. If we assume that this alternative definition was operative 139 in the study for at least some of the women and their reported LMP dates, then the classifications made in the 140 reported secondary analyses were wrong and the conclusion that L H oophorectomy surgery gives better outcomes 141 was grounded in mis-classifications [4]. If the conclusions from the new adjuvant (reported here) and metastatic 142 c) Menstrual cycle hormonal biology which may explain the new surgical oophorectomy timing findings What 143 144 biological explanation is consistent with the summarized data that prolonged follicular phase patients derive minimal benefit from surgical oophorectomy plus tamoxifen treatments? To begin, it is important to note that 145 typical human levels of progesterone are < 1 nanogram (ng) to about 20 ng/ml, while levels of estradiol are 146 50-200 picograms (pg)/ml. Thus, a typical luteal phase level of progesterone of 10 ng/ml is 50-fold greater than 147 a typical estradiol level of 200pg/ml. 148

When ovulation is delayed, there are sustained high estradiol levels for as many as 14 days or more. Indeed, 149 in our data, the mean estradiol levels on the day of surgery were higher in the prolonged follicular phase (or L H 150 unconfirmed) group of patients than in the confirmed follicular patients. In the surgical opphorectomy situation, 151 no progesterone "rescue" follows. In normal follicular phase, estradiol exposure is short, and in normal luteal 152 phase exposure to some duration of progesterone "rescue' occurs before the oophorectomies. In anovulatory 153 patients, the high and prolonged estradiol levels stimulate growth of micrometastases as the last hormonal signal 154 155 that these lesions receive. When it is done during the follicular phase of a cycle, oophorectomy appears to 156 send a strong anti-growth signal. A flare of the metastatic disease is often seen about 7-10 days after starting 157 the treatment. This kind of flaremay be what is occurring with follicular phase oophorectomy. In a normal luteal phase, oophorectomy may have relatively small acute effects because of the last signals, which are high 158 progesterone level-mediated. The data from our two trials collectively are showing extraordinarily limited effects 159 (in the sense of limited/no benefit from oophorectomies plus tamoxifen) in designated prolonged follicular phase-160 low progesterone patients from limited-time hormonal differences, while showing strong effects when this surgery 161 is done in usual follicular or high progesterone luteal phases. 162

163 8 Medical

164 V.

165 9 Conclusions

The potential greater efficacy with timing in the menstrual cycle of the surgical oophorectomy would make 166 this treatment combined with tamoxifen, already the first global option adjuvant treatment based on efficacy, 167 practicality and cost-efficacy, an even more compelling therapy [2,3]. A practical interpretation is that acting on 168 this observation and performing surgical oophorectomies whenever possible in hormonally confirmed follicular or 169 luteal phases appears very unlikely to be harmful in terms of efficacy. Were surgical oophorectomy plus tamoxifen 170 adjuvant therapy widely promoted and applied across the world, a reasonable estimate is that 100,000 women a 171 year would be saved, women who otherwise would get little or no effective adjuvant treatment (12). Were timed 172 surgical oophorectomy widely promoted and applied as hostpersonalized therapy, an additional 20,000 women 173 per year might be saved. 174

If a conservative position is taken with regard to these timing data, that the case that women in prolonged follicular phase with low progesterone levels benefit little from oophorectomy done at this time, has but limited support, then the rational approach is to do a clinical trial of timed SO+T (excluding prolonged follicular phase confirmed women) vs. GnRH/LHRH +T (or aromatase inhibitor). With provision of the drugs, this would not be a difficult trial to do, certainly with low-and middle-income county participation.

180 10 Declarations

¹⁸¹ 11 Ethics approval and consent to participate

The data reported in this manuscript have come from previously approved clinical trials. The approvals have been both in the home countries of the patients and in the United States.

¹⁸⁴ 12 Consent for publication

185 With this submission the sole author implicitly provides consent to publish.

13 d) Other data which bear on the new hypothesis/ interpre tations

There are five observations which validate our findings because they are consistent with our observation of 188 limited benefit from prolonged follicular phase patient-surgical oophorectomy. First, there are immediate and 189 severe vasomotor symptoms in women following surgical oophorectomy. Second, men with metastatic prostate 190 cancer have immediate responses with decreases in bone pain following orchiectomy. Third, Badwe et al. found 191 that short-term adjuvant, parenteral peri-operative progesterone, which was associated with better outcomes in 192 axillary node positive patients [8]. These results are consistent with our observation of absence of benefit with 193 low-progesterone prolonged follicular phase patients. Four, the peaks of hazards for recurrence of breast cancer 194 at 2-3 years post diagnosis and treatment have most strongly been related to peri-operative changes. 195

Baum et al. suggested that minor peri-operative changes can lead to major long-term effects [9,10]. Finally, other perioperative conditions of limited duration have been suggested to have major longer-term impacts [11]. Scheduled surgeries were assigned to be in mid-luteal phase by history. For these patients, by history, 96% of surgeries were done in luteal phase. For these patients, 66% had surgeries by history in follicular phase These percentages make clear the rationale for the secondary analyses based on the better menstrual cycle phase status of study patients using day of surgery progesterone levels.

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206 .2 Availability of data and material

207 The primary study data and files are available. ClinicalTrials.govnumbers, NCT 00201851 and NCT00293540

208 .3 Competing interests

209 The author reports no conflicts of interest.

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213 .5 Authors' contributions

- 214 The sole author is responsible for all parts of this report.
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