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# Predicting the cumulative chance of live birth over multiple complete cycles of in vitro fertilisation: an external validation study

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### 26 ABSTRACT

27

28 Study question

Are the published pre-treatment and post-treatment McLernon models, predicting cumulative live birth
 rates (LBR) over multiple complete IVF cycles, valid in a different context?

31

32 *Summary answer* 

With minor recalibration of the pre-treatment model, both McLernon models accurately predict
 cumulative LBR in a different geographical context and a more recent time period.

35

36 What is known already

37 Previous IVF prediction models have estimated the chance of a live birth after a single fresh embryo 38 transfer, thereby excluding the important contribution of embryo cryopreservation and subsequent IVF 39 cycles to cumulative LBR. In contrast, the recently developed McLernon models predict the cumulative 40 chance of a live birth over multiple complete IVF cycles at two certain time points: a) before initiating 41 treatment using baseline characteristics (pre-treatment model) and b) after the first IVF cycle adding 42 treatment related information to update predictions (post-treatment model). Before implementation of 43 these models in clinical practice, their predictive performance needs to be validated in an independent 44 cohort.

45

46 Study design, size, duration

External validation study in an independent prospective cohort of 1515 Dutch women who participated in
the OPTIMIST study (NTR2657) and underwent their first IVF treatment between 2011 and 2014.
Participants underwent a total of 2881 complete treatment cycles, with a complete cycle defined as all
fresh and frozen thawed embryo transfers resulting from one episode of ovarian stimulation. The follow

up duration was 18 months after inclusion, and the primary outcome was ongoing pregnancy leading tolive birth.

- 53
- 54 *Participants/materials, setting, methods*

55 Model performance was externally validated up to three complete treatment cycles, using the linear 56 predictor as described by McLernon et al. to calculate the probability of live birth. Discrimination was 57 expressed by the c-statistic and calibration was depicted graphically in a calibration plot. In contrast to the 58 original model development cohort, anti-Müllerian hormone (AMH), antral follicle count (AFC) and body 59 weight were available in the OPTIMIST cohort, and evaluated as potential additional predictors for model 50 improvement.

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#### 62 *Main results and the role of chance*

63 Applying the McLernon models to the OPTIMIST cohort, the c-statistic of the *pre-treatment model* was 64 0.62 (95% confidence interval (CI) 0.59-0.64) and of the post-treatment model 0.71 (95% CI 0.69-0.74). 65 The calibration plot of the *pre-treatment model* indicated slight overestimation of the cumulative LBR. To 66 improve calibration, the *pre-treatment model* was recalibrated by subtracting 0.35 from the intercept. The post-treatment model calibration plot revealed accurate cumulative LBR predictions. After addition of 67 68 AMH, AFC and body weight to the McLernon models, the c-statistic of the updated pre-treatment model 69 improved slightly to 0.66 (95% CI 0.64-0.68), and of the updated post-treatment model remained at the 70 previous level of 0.71 (95% CI 0.69-0.73).

Using the *recalibrated pre-treatment model*, a woman aged 30 years with two years of primary infertility who starts ICSI treatment for male factor infertility has a chance of 40% of a live birth from the first complete cycle, increasing to 72% over three complete cycles. If this woman weighs 70 kilograms, has an AMH of 1.5 ng/mL and an AFC of 10 measured at the beginning of her treatment, the *updated pretreatment model* revises the estimated chance of a live birth to 30% in the first complete cycle and 59% over three complete cycles. If this woman then has 5 retrieved oocytes, no embryos cryopreserved and a

single fresh cleavage stage embryo transfer in her first ICSI cycle, the *post-treatment model* estimates the
chances of a live birth at 28% and 58%, respectively.

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# 80 *Limitations, reasons for caution*

Two randomised controlled trials (RCT) evaluating the effectiveness of gonadotropin dose individualisation on basis of the AFC were nested within the OPTIMIST study. The strict dosing regimens, the RCT in- and exclusion criteria and the limited follow up time of 18 months might have influenced model performance in this independent cohort. Also, consistent with the original model development study, external validation was performed using the optimistic assumption that the cumulative LBR in couples who discontinue treatment without a live birth would have been equal to that of those who continue treatment.

88

## 89 Wider implications of the findings

After national recalibration to account for geographical differences in IVF/ICSI treatment, the McLernon prediction models can be introduced as new counselling tools in clinical practice to inform patients and to complement clinical reasoning. These models are the first to offer an objective and personalised estimate of the cumulative probability of live birth over multiple complete IVF cycles.

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106	
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108	Prediction model, external validation, live birth, IVF/ICSI, infertility, cumulative live birth, personalised,
109	counselling, prognostic research

### 110 Introduction

111 Infertility is defined as the failure to conceive within 12 months of regular unprotected intercourse, and 112 affects approximately one in six couples (Oakley et al., 2008; Zegers-Hochschild et al., 2017). The 113 majority of infertile couples seek fertility care, and many of those with prolonged unresolved infertility 114 will be treated with ART regardless of cause (Boivin et al., 2007; Datta et al., 2016). IVF and ICSI are 115 both widely used techniques for couples with infertility. Globally more than 1.6 million annual cycles of 116 IVF/ICSI are performed and while success rates have increased over time (Dyer et al., 2016; McLernon et 117 al., 2016), this treatment is still not effective for all infertile couples, with live birth rates (LBR) at around 118 25-30% per treatment cycle (Malizia et al., 2009; McLernon et al., 2016; de Neubourg et al., 2016). Since 119 IVF/ICSI is expensive and carries several risks, the probability of a live born child should be weighed 120 against the risks and costs of this treatment.

121 Several prognostic models have been developed to objectively estimate the probability of a live birth after 122 IVF/ICSI treatment (Leushuis et al., 2009; van Loendersloot et al., 2014). It is known that prediction 123 models often perform optimistically in their development sample, even after correction by internal 124 validation. This is caused by overfitting, which occurs when the model corresponds too closely to the 125 development data due to the inclusion of too many predictors (Moons, Kengne, Woodward, et al., 2012). 126 External validation in an independent cohort of women is thus essential to examine the performance and 127 generalisability of the prediction model (Altman et al., 2009; Harrell et al., 1996). Unfortunately, most of 128 the currently available models that predict the chance of a live birth after IVF/ICSI treatment have never 129 been externally validated (Leushuis et al., 2009; van Loendersloot et al., 2014). Also, the majority of 130 these models predict the probability of a live birth after a single fresh embryo transfer, excluding the 131 important contribution of embryo cryopreservation and subsequent treatment cycles to LBR. This limits 132 their potential as counselling tools for couples and clinicians, especially considering the increased use and 133 improved techniques of embryo cryopreservation and frozen thawed embryo transfer cycles in recent 134 years (Wong et al., 2014).

135 Three of the largest model development studies for prediction of live birth after IVF and/or ICSI 136 treatment used data from the Human Fertilisation and Embryology Authority (HFEA) database in the UK 137 (McLernon et al., 2016; Nelson and Lawlor, 2011; Templeton et al., 1996). Treatment and outcome data 138 from all licenced fertility clinics within the UK have been recorded in this database since 1992. The two 139 models developed by Templeton et al. and Nelson et al. were both externally validated, and their 140 predictive performance was compared to one another in several studies (Arvis et al., 2012; van 141 Loendersloot et al., 2011; Smeenk et al., 2000; Smith et al., 2015; te Velde et al., 2014). Although these 142 models have been recommended in previous studies and used internationally to predict live birth after 143 IVF and ICSI (Leushuis et al., 2009; Smith et al., 2015; te Velde et al., 2014), neither model predicts 144 cumulative LBR over multiple IVF/ICSI treatment cycles including frozen thawed embryo transfer 145 cycles.

146 Recently, a new model was developed by McLernon et al. using the HFEA database (McLernon et al., 147 2016). This model is the first to provide an individualised estimate of the cumulative chance of a live 148 birth over multiple complete cycles of IVF/ICSI, with a complete cycle defined as all fresh and frozen 149 thawed embryo transfers resulting from one episode of ovarian stimulation. For model development, data 150 from 113 873 women and 184 269 complete cycles between 1999 and 2009 were used. Internal validation 151 of the model showed promising results, however evaluation of the predictive performance of the model in 152 a different geographical context using more contemporary data has yet to be performed. Additionally, a 153 number of potential key predictors, such as measures for ovarian reserve and female body weight, were 154 unavailable in the HFEA database and could not be included in the original model (McLernon et al., 155 2016).

156 The main objective of the current study was therefore to perform geographical and temporal validation of 157 the new HFEA model by using recent data from a different country. We also wanted to determine whether 158 inclusion of additional parameters, such as female body weight and ovarian reserve test results i.e. antral

- 159 follicle count (AFC) and anti-Müllerian hormone (AMH), could improve the predictive performance of
- the model.

#### 161 Materials and methods

#### 162 Data sources

163 External validation was performed on data from the OPTIMIST study (van Tilborg, Oudshoorn, et al., 164 2017). This multicentre prospective cohort study included 1515 women from 25 infertility centres in the 165 Netherlands between May 2011 and May 2014. Participants were younger than 44 years of age, had 166 regular menstrual cycles and no significant uterine or ovarian abnormalities on transvaginal ultrasound. 167 Women with polycystic ovarian syndrome, metabolic or endocrine abnormalities or undergoing oocyte 168 donation were excluded. All participants were included before their first IVF/ICSI cycle, or the first cycle 169 after a previous live birth. The primary outcome was ongoing pregnancy, achieved within 18 months of 170 follow up, and resulting in live birth. Ethical approval for the OPTIMIST study was obtained from the 171 Institutional Review Board of the University Medical Centre Utrecht (MEC 10-273), and all participants 172 provided written informed consent. A more detailed description of study procedures and results were 173 reported previously (Oudshoorn et al., 2017; van Tilborg et al., 2012; van Tilborg, Oudshoorn, et al., 174 2017; van Tilborg, Torrance, et al., 2017).

## 175 McLernon model

The McLernon model consists of two clinical prediction models to estimate the individualised cumulative chance of a live birth over a maximum of six complete treatment cycles. **Before initiating treatment**, the *pre-treatment model* predicts the probability of a live birth from both fresh and frozen thawed embryo transfers based on couple characteristics and the use of IVF or ICSI. Included predictors are: female age (years), duration of infertility (years), previous pregnancy, causes of infertility (tubal factor, anovulation, male factor, unexplained infertility), type of treatment (IVF or ICSI) and treatment year (see Supplementary Text 1).

183 After the first fresh treatment cycle, treatment specific characteristics from this cycle are added in the 184 *post-treatment model* to update the predicted probability. Added predictors are: number of oocytes,

185 cryopreservation of embryos, and the number and stage of embryos at the first fresh embryo transfer 186 (single, double or triple embryo transfer; blastocyst or cleavage stage). All causes of infertility are 187 excluded as predictors in the post-treatment model, except for tubal factor (see Supplementary Text 2). 188 For women with zero oocytes collected in the first cycle, a separate post-treatment model is available.

To predict the probability of a live birth in the *i*th cycle, assuming no live birth occurred in the previous cycle(s), complete cycle number is included in both models as a discrete time variable. A complete cycle includes all fresh and frozen thawed embryo transfers resulting from one episode of ovarian stimulation. With the predicted probability of a live birth per subsequent complete cycle, the cumulative probability of a live birth can be calculated up to six complete cycles (see Supplementary Text 1 and 2).

### 194 *Statistical analysis*

Nine predictor variables had missing values (Table I). The proportion of missing values was low (< 2.5%), except for AMH (11.2%). During the OPTIMIST study, blood sampling was performed on the day of randomisation. Logistic issues prevented blood sampling in some cases, thus compromising the ability to undertake post-hoc measurements of AMH in the total population. As the reasons for missing values were considered to be unrelated to the AMH value itself or the measurement, these were defined as missing (completely) at random.</p>

201 Multiple imputation was applied for predictors with missing values in the OPTIMIST database (Sterne et 202 al., 2009). In this process 10 imputed datasets were created using a multivariate imputation by chained 203 equations (MICE) algorithm (van Buuren and Groothuis-Oudshoorn, 2011). Predicted probabilities for a 204 live birth were calculated on each imputed dataset, using the predictors and parameter-estimates of both 205 the pre-treatment model as well as the post-treatment model as described by McLernon et al 2016 206 (McLernon et al., 2016). In accordance with the original models, the variables female age, treatment year 207 and number of oocytes were treated with restricted cubic splines in the validation process. The separate 208 post-treatment model for women with zero oocytes collected in the first treatment cycle was not validated

in this study, as the number of women for this analysis was too low in the OPTIMIST database. Cumulative probabilities were calculated up to three complete IVF/ICSI cycles, as most couples in the Netherlands only have three treatment cycles due to the current reimbursement policy. Also, the OPTIMIST follow up period was 18 months, reducing the number of women with more than three treatment cycles. The validation process was performed ten times on each of the imputed datasets and separate results were pooled using Rubin's rules (Rubin, 2004).

215 The predictive performance of the McLernon models was evaluated in terms of discrimination and 216 calibration. Discrimination quantifies the ability of a model to correctly differentiate between subjects 217 with an event and subjects without an event (Moons, Kengne, Woodward, et al., 2012). In the context of 218 fertility treatment, it is the ability of the models to distinguish between women with a live birth and 219 women without a live birth after IVF/ICSI treatment. It is expressed by the c-statistic or the area under the 220 receiver operating curve (AUROC), which ranges between 0.5 and 1. A c-statistic of 1 indicates perfect 221 discrimination, whereas a c-statistic of 0.5 represents a model with no discrimination at all. In this study, 222 the c-statistic (and 95% CI) was calculated using the method suggested by Harrell et al., (Harrell et al., 223 1996).

224 Calibration describes the degree of agreement between predicted probabilities and observed outcomes 225 (Moons, Kengne, Woodward, et al., 2012), in this context the predicted probability of a live birth and the 226 observed LBR. Calibration can be assessed graphically by forming subgroups of patients determined by 227 ranges of predicted probabilities, and then plotting the observed proportion of events against the mean 228 predicted probability within these subgroups. When perfect calibration is present, the plot shows a 229 diagonal line with a slope of one and an intercept of zero. In the current study, five equal subgroups of 230 patients were formed. This was based on the sample size of the OPTIMIST cohort and the related 231 precision of the point estimates in the calibration plot. Within these subgroups, the Kaplan Meier 232 estimates of the observed cumulative LBR over three complete treatment cycles were plotted against the 233 mean predicted probability of cumulative live birth. A smoothed line was then added in this plot using the

proportional hazard regression approach described by Harrell et al (Harrell *et al.*, 1996). In addition to this, a systematic difference in the predicted and observed LBR was assessed by using calibration-in-thelarge (Steyerberg, 2009), and the intercept of the prediction models was adjusted in case a systematic over- or underestimation was present.

238 *Updating the models* 

Following the external validation of the models, the additional value of updating the McLernon models with pre-specified new biomarkers was evaluated. AMH (ng/mL), AFC (2-10 mm) and body weight (kg) were added to the pre-treatment and post-treatment model in a multivariable logistic regression analysis, in which the linear predictor of the McLernon model was entered as a fixed variable. The final model was established using a manual backward selection process. Predictors were eliminated from the model according to the Akaike Information Criterion (AIC) (Akaike, 1974).

245 The predictive performance of the new updated models was evaluated by calculating the c-statistic (and 246 95% CI). To assess for overfitting, internal validation was performed by bootstrapping (Steverberg, 247 2009). Two hundred bootstrap samples, all of which were of the same size as the original validation 248 sample, were created by random sampling with replacement (Harrell, 2001; Steyerberg, 2009). In each 249 bootstrap sample, a new model was fitted with the same predictors as the updated models. The c-statistic 250 was calculated for each of the 200 sample derived models, in both the bootstrap sample as well as the 251 original validation cohort. The difference between these two c-statistics was calculated for each of the 200 252 sample derived models, and averaged to give the optimism estimate. This was subtracted from the 253 original c-statistic to obtain the optimism corrected c-statistic for the updated models.

All statistical analyses were performed using R for Windows (version 3.3.2; R Foundation for Statistical
Computing, Vienna, Austria).

#### 256 **Results**

257 Of the 1515 women included in the OPTIMIST study, four were excluded in the current study as they 258 never started IVF/ICSI treatment. A total of 2881 IVF/ICSI cycles were performed over a period of 18 259 months of follow up. Table I shows the patient and first cycle treatment characteristics of the OPTIMIST 260 cohort (validation sample) and the HFEA cohort (development sample). Women included in the 261 validation sample were about the same age as women in the development sample, but had a shorter 262 average duration of infertility. The causes of infertility showed a similar distribution across both samples, 263 with the exception of anovulation which rendered women ineligible for the OPTIMIST study. The 264 treatment characteristics showed that embryo cryopreservation was more frequently performed after the 265 first IVF/ICSI cycle in the validation sample and that these women most often had a cleavage stage single 266 embryo transfer in the first fresh cycle, whereas women in the development sample most often had a 267 cleavage stage double embryo transfer. No formal assessment was performed for the differences and 268 similarities between the cohorts, as a description rather than a p-value is considered to be useful for 269 interpretation of the models' performance in this external validation study.

270 The flowchart in Figure 1 shows the number of women in the OPTIMIST and HFEA cohorts who started 271 a treatment cycle, had a live birth or discontinued treatment without having a live birth. The LBR per 272 cycle was similar in both cohorts for the first, second and fourth treatment cycle. In the third cycle the 273 LBR was slightly higher in the OPTIMIST cohort compared to the HFEA cohort. As few women in the 274 OPTIMIST cohort received a fifth or sixth cycle, LBR in these cycles could not be compared. The 275 proportion of women without a live birth that continued treatment was higher after the first and second 276 cycle in the OPTIMIST cohort as compared to the HFEA cohort. After the third cycle, the proportion 277 continuing treatment in the OPTIMIST cohort decreased, while it remained constant in the HFEA cohort. 278 At the end of follow up, 52% of the women in the OPTIMIST study had a treatment related live birth. The 279 overall LBR of the HFEA cohort was 43% over six complete IVF/ICSI cycles.

As mentioned previously, external validation of the McLernon models was performed up to three complete treatment cycles, and therefore the fourth, fifth and sixth complete treatment cycle in the OPTIMIST dataset (n=102 complete treatment cycles, n= 15 live births) were excluded from further analysis. Also, for the post-treatment model validation, women with zero oocytes collected in the first treatment cycle were excluded (n= 226 women, n = 526 complete treatment cycles, n= 82 live births) as a separate model was developed for this group of women by McLernon et al (McLernon *et al.*, 2016). Due to the small numbers, this separate model could not be validated in this study.

# 287 Discrimination and calibration

In the validation sample, the pooled c-statistic for the pre-treatment model was 0.62 (95% CI 0.59-0.64) and for the post-treatment model 0.71 (95% CI 0.69-0.74). Figure 2a and 3 show the calibration plots for both original models, depicting the correlation between the observed and predicted cumulative LBR. The pre-treatment calibration plot had an intercept of -0.23 (95% CI -0.36- -0.10) and a slope of 0.98 (95% CI 0.69-1.27), and the post-treatment calibration plot had an intercept of -0.01 (95% CI -0.12-0.11) and a slope of 0.97 (95% CI 0.77-1.19).

294 The pre-treatment model systematically overestimated the cumulative LBR over three complete cycles for 295 women in the validation sample. This is shown by a calibration curve with most of the confidence 296 intervals under the reference line (Figure 2a), indicating significantly higher predicted probabilities than 297 observed LBR. The calibration-in-the-large analysis confirmed this systematic overestimation with an 298 intercept of -0.35. To improve calibration, the pre-treatment model was thus adjusted by subtracting 0.35 299 from the intercept of the original linear predictor, which decreased the predicted odds of a live birth by a 300 factor of 1.42 (see Supplementary Text 3). The calibration plot of the recalibrated pre-treatment model 301 showed improved accuracy of the predictions, with all confidence intervals overlapping the reference line 302 (Figure 2b). In contrast to the pre-treatment model, the post-treatment model correctly estimated the cumulative LBR in the validation sample, as is shown by a calibration plot with confidence intervals
overlapping the reference line indicating no significant over- or underestimation (Figure 3).

## 305 Updating of the models

306 Addition of the biomarkers AMH, AFC and body weight to the pre-treatment and post-treatment model in 307 a multivariable regression analysis resulted in two new updated models. The updated pre-treatment model 308 included all three biomarkers as additional predictors for live birth. Since the relationship between both 309 AMH and AFC with the probability of live birth was non-linear, these predictors were included using 310 restricted cubic splines (see Supplementary Figure 1). The updated post-treatment model included only 311 AFC and AMH as additional predictors for live birth, of which AFC was modelled by using restricted 312 cubic splines (see Supplementary Figure 2). After internal validation of the updated models by bootstrapping, the updated pre-treatment model had a corrected c-statistic of 0.66 (95% CI 0.64-0.68) and 313 314 the updated post-treatment model had a corrected c-statistic of 0.71 (95% CI 0.69-0.73). The addition of 315 AFC, AMH and body weight thus resulted in a slight improvement of the discriminatory capacity of the 316 pre-treatment model, while addition of AFC and AMH had no beneficial effect on the discriminative 317 performance of the post-treatment model.

#### 318 *Examples of model predictions*

319 Figures 4, 5 and 6 show examples of model predictions as illustration for clinical application. Figure 4 320 presents predictions of the *recalibrated pre-treatment model* for couples with primary infertility caused 321 by a male factor. Cumulative probabilities of live birth are calculated up to three complete ICSI cycles, 322 and are differentiated by female age (30 or 40 years) and duration of infertility (2 years or 5 years). As is 323 shown in figure 4, age is the most important predictor in the pre-treatment model. A 30-year-old woman 324 with 2 years of infertility has a predicted probability of a live birth of 0.40 in the first ICSI cycle, 325 increasing to 0.72 over three complete cycles. For a 40-year-old woman with 2 years of infertility, these 326 probabilities are 0.15 and 0.32 respectively.

327 Figure 5 shows predictions of the *updated pre-treatment model*, with AMH, AFC and body weight as new 328 predictors in the model. Predictions are presented for couples with two years of primary infertility caused 329 by a male factor, and differentiation is based on female age (30 or 40 years), AMH (2.0 or 0.5 ng/mL) and 330 AFC (15 or 7). In all scenarios the female body weight is 70 kilograms. A 30-year-old woman with an 331 average ovarian reserve at the start of her first treatment - indicated by an AMH of 2.0 ng/mL and an 332 AFC of 15 - has a predicted probability of a live birth of 0.37 in the first cycle and 0.69 over three cycles333 (0.17 and 0.37 for a 40-year-old woman). If this woman has a reduced ovarian reserve – indicated by an 334 AMH of 0.5 ng/mL and an AFC of 7 – the predicted probabilities decrease to 0.19 and 0.42, respectively 335 (0.08 and 0.18 for a 40-year-old woman).

336 Figure 6 shows predictions of the *post-treatment model*, which revises the predicted probabilities of the 337 pre-treatment models by adding information of the first treatment cycle. Predictions are calculated for 338 women with two years of primary, non-tubal infertility and are differentiated by female age (30 or 40 339 years), number of oocytes (10 or 5) and embryo cryopreservation (yes or no). In all scenarios the woman 340 received a cleavage stage single embryo transfer. The predicted probabilities of a live birth for women 341 with a favourable prognosis – aged 30-years, 10 oocytes retrieved and cryopreserved embryos – is 0.49 in 342 the first ICSI cycle, increasing to 0.83 over 3 complete cycles. In contrast, for women with a poorer 343 prognosis – aged 40 years, 5 oocytes retrieved and no embryos cryopreserved – the predicted probabilities 344 are 0.11 and 0.26, respectively.

#### 345 Discussion

#### 346 *Main findings*

This external validation study of the McLernon pre-treatment and post-treatment model found that, after minor recalibration of the intercept of the pre-treatment model, both models accurately predict the cumulative probability of live birth up to three complete IVF/ICSI cycles in a more contemporary cohort in another country. The discriminatory capacity of the pre-treatment model in an external cohort was limited, whereas the post-treatment model had a fair ability to discriminate between couples with and without a live birth after treatment.

## 353 Strengths

This study focuses on the external validation of an IVF prediction model, which is an essential but frequently overlooked step before implementation of a prediction model in clinical practice (Altman *et al.*, 2009). In contrast to redeveloping new models for the same outcome, external validation and updating of existing models prevents the loss of scientific information by combining the information captured in the original model with information of a new patient cohort (Moons, Kengne, Grobbee, *et al.*, 2012).

Embryo cryopreservation has become an important part of IVF/ICSI treatment, and most couples have more than just one complete treatment cycle (Wong *et al.*, 2014). Unlike previous prediction models (Leushuis *et al.*, 2009; van Loendersloot *et al.*, 2014), the McLernon models provide a more useful estimate of cumulative treatment success. As such, the validation of these models represents a significant step forward in creating a clinically useful tool to manage expectations and to inform decision making around IVF.

This study benefits from the prospective design of the OPTIMIST study, which has ensured reliable data collection, with relatively low numbers of missing values and a low risk of selection bias. The multicentre design resulted in a highly representable cohort for Dutch fertility care. And although it is known that the

368 IVF/ICSI success rates vary between fertility centres, the inclusion of multiple centres will increase the 369 generalisability and applicability of the external validation of the McLernon models within the 370 Netherlands.

Furthermore, the external validation was performed on data collected in a recent time period (2011-2014). Due to changing patient populations, new treatment protocols, improving technologies and increasing success rates over time, prediction models in reproduction medicine have no static form and should be regularly updated to optimally reflect the latest circumstances in which they are used (Altman *et al.*, 2009). As the McLernon models were developed on data collected between 1999 and 2009, data of the more recently performed OPTIMIST study were helpful to investigate if model performance was still accurate in current practice.

#### 378 Weaknesses

This study has a number of limitations. First, the external validation involved data from a prospective cohort study within which two randomised controlled trials were embedded evaluating the effectiveness of individualised doses of gonadotropins based on AFC. Strict dosing regimens might have affected some treatment outcomes, such as cancellation rates and number of oocytes, thus influencing the predictive capacity of the models in the validation sample. However, as the OPTIMIST study found no difference between the dosing regimens on cumulative live birth rates, the impact on model performance is likely to be minimal.

Second, the OPTIMIST study used strict eligibility criteria. Therefore, the validation sample does not fully represent the diversity of the patient population initiating IVF/ICSI treatment in the Netherlands. As none of the women in the validation sample were anovulatory, external validation of the models was only performed for an ovulatory population. This limits the generalisability of the models to some extent, as the original McLernon models were developed in a population which also included anovulatory women. Also, it could have had some impact on model performance. However, since anovulation had only a small

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392 predictive value in the pre-treatment model, and the majority of couples underwent IVF/ICSI for other393 indications, a large impact on model performance is unlikely.

Third, the OPTIMIST study had a follow up period of 18 months, leading to small numbers of women with more than three complete treatment cycles. Model performance could therefore only be reliably validated up to three complete cycles. However, most couples in the Netherlands complete a maximum of three treatment cycles which is partly due to the national reimbursement policy, but also by the high rates of embryo cryopreservation, increasing the number of embryo transfers and LBR per cycle. Therefore, model validation up to three complete cycles has particular clinical relevance for current Dutch fertility care.

401 Last, the original McLernon prediction models were developed on linked cycle data, which were then 402 used to estimate cumulative pregnancy chances. Therefore, these models used the optimistic assumption 403 that the cumulative LBR in couples who discontinue IVF treatment without a live birth would have been 404 equal to that of couples who continue further treatment cycles, after correction of predictor effects. This 405 assumption tends to lead to overestimation of the cumulative LBR, as women with a low prognosis of 406 achieving a live birth are generally more likely to discontinue treatment (Brandes et al., 2009; Olivius et 407 al., 2004). Since the reasons for treatment withdrawal were unknown in the current external validation 408 study, a similar method was used that probably resulted in some degree of overestimation of the 409 cumulative LBR in the validation cohort. However, as the original McLernon models were developed 410 with this approach, and the predictions for cumulative LBR over multiple complete cycles were 411 considered to be clinically more relevant than per cycle predictions, we feel that the current method is the 412 best option for the external validation of the McLernon models.

### 413 *Explanation of findings*

The discriminatory capacity of the pre-treatment model was markedly lower in the validation sample than in the development sample. In the development study, a c-statistic of 0.73 (95% CI 0.72-0.74) was

416 reported, whereas the present study found a c-statistic of 0.62 (95% CI 0.59-0.64). For the post-treatment 417 model, the discriminatory performance in the validation sample was comparable to that in the 418 development sample, with a c-statistic of 0.71 (95% CI 0.69-0.74) and 0.72 (95% CI 0.71-0.73) 419 respectively (McLernon et al., 2016). As it is known that prediction models tend to perform too 420 optimistically in the development dataset due to overfitting, some reduction in model performance is to be 421 expected during external validation due to the differences between samples (Altman et al., 2009; Moons, 422 Kengne, Woodward, et al., 2012). This, to some extent, also explains the lower overall performance of 423 the pre-treatment model. The comparable performance of the post-treatment model in both samples 424 indicates that the treatment related variables that were added to this model (number of oocytes, 425 cryopreservation of embryos, and the number and stage of embryos) are important predictors for live birth 426 after treatment.

427 Other than the influence of overfitting, some key differences between the Dutch and UK healthcare 428 systems may also have affected the models' performance in this external validation study. An important 429 factor is the reimbursement policy for fertility treatment. All Dutch infertile couples are insured for a 430 minimum of three complete IVF/ICSI cycles. In contrast, most couples in the UK receive no standard 431 funding for ART (Berg Brigham et al., 2013). Since IVF/ICSI treatment is expensive, this induces 432 discrepancies in the patient population initiating and continuing treatment between the two study samples 433 (Rajkhowa et al., 2006). As can be seen in the baseline table (Table I) and flowchart (Figure 1), couples 434 in the UK had a longer average duration of infertility before starting treatment and were more likely to 435 discontinue treatment after the first and second cycles than couples in the Netherlands. Also, the decrease 436 in LBR is more evident in the UK than in the Netherlands over the first three cycles, which suggests that 437 differences exist in both reasons for discontinuation as well as prognostic profiles of women 438 discontinuing treatment in the two countries. These phenomena are, in part, financially driven, and could 439 partially explain the difference in predictive ability of the UK models in the Dutch cohort.

440 Furthermore, despite the fact that the infertility guidelines of both countries include similar approaches 441 for treatment of infertile couples, there are important variations in treatment characteristics between the 442 two study samples (Dutch Society of Obstetrics and Gynaecology (NVOG), 2010; National Institute for 443 Health and Care Excellence (NICE), 2013). Some of these differences are mainly due to changes in 444 clinical practice over time. As is shown by the baseline table (Table 1), women in the more recent Dutch 445 cohort (2011-2014) generally had a single embryo transfer in their first fresh treatment cycle, whereas 446 women in the earlier UK cohort (1999-2009) most often had a double embryo transfer. Also, embryo 447 cryopreservation was performed in over half of the Dutch women as compared to only a guarter of the 448 women in the UK. Other differences are explained by variation in treatment protocols between 449 geographic locations. For one, no blastocyst stage embryos transfers were performed in the Netherlands in 450 contrast to the proportion of blastocyst stage embryo transfers in the UK of more than 10%. Also, Dutch 451 women more frequently had no embryo available for transfer after their first treatment cycle, which is 452 most likely caused by strict cancellation criteria particularly for hyper response. These differences in 453 treatment characteristics suggest that the development sample does not fully reflect clinical practice in a 454 more recent time period and in a different geographic context. As cumulative LBR are substantially 455 affected by the variation in treatment characteristics (Glujovsky et al., 2016; Pandian et al., 2013; Wong 456 et al., 2014), this could explain part of the different performance of the pre-treatment model in the 457 validation sample. The stable performance of the post-treatment model, which includes embryo stage and 458 embryo cryopreservation as important predictors, seems to confirm the impact of the variation in these 459 variables on model performance.

The addition of measures of ovarian reserve, i.e. AMH and AFC, and body weight to the McLernon prediction models revealed only a marginal improvement of model performance in the OPTIMIST dataset. The additional value of these tests can therefore be questioned, especially in view of the extra costs and physical burden on the patient. Female age is one of the most important predictors in the McLernon models (McLernon *et al.*, 2016). As female age is correlated with the ovarian reserve, adding

AMH and AFC provides limited new information to the prediction models. This is in line with previous studies that showed that ovarian reserve tests have no added value to the use of female age alone in the prediction of ongoing pregnancy after treatment (Broer *et al.*, 2013). Other potential predictors for live birth, such as ethnicity, smoking status and alcohol intake, were not included in this update of the McLernon model (Dhillon *et al.*, 2015; Rossi *et al.*, 2011; Waylen *et al.*, 2009). The additional value of these variables for model performance was considered uncertain, as the reporting is remarkably subjective and/or often incomplete (Liber and Warner, 2018; Stockwell *et al.*, 2016).

### 472 *Clinical implications*

473 Discrimination and calibration have been recognized as measures to evaluate the performance of 474 prediction models (Altman et al., 2009; Steyerberg, 2009). However, the discriminative ability at the 475 binary level of most prediction models in reproductive medicine, as expressed by the c-statistic, is 476 considerably low (Leushuis et al., 2009). As at the moment of prediction the outcome of pregnancy has 477 not yet occurred, the c-statistic is determined using the calculated probability of pregnancy. The 478 maximum value of the c-statistic depends on the variability of these calculated probabilities in the 479 infertile population. Since infertility is a complex and multifactorial health problem and due to the 480 absence of strong predictors for live birth – particularly pre-treatment –, the probability distribution in 481 infertile couples that have a live birth has a considerable overlap with the distribution of those without a 482 live birth. Therefore the maximum c-statistic can be expected to be low (Cook, 2007; Coppus et al., 483 2009), as is seen in the external validation of the pre-treatment model. However, this does not necessarily 484 imply that such prediction models have limited use in clinical practice. Models with reliable predictions 485 and a clinically useful distribution of probabilities for achieving a live birth, as assessed by calibration, 486 can still support patients and clinicians in clinical decision making around infertility treatment (Coppus et 487 al., 2009).

488 As the calibration plots of both the recalibrated pre-treatment model and the post-treatment model 489 indicate accurate predictions with a useful range of prognoses, these models can be used within the 490 Netherlands as counselling tools to complement clinical reasoning at two certain time points. Before 491 initiating treatment, the recalibrated pre-treatment model offers couples and clinicians a personalised and 492 objective estimate of success over multiple complete treatment cycles. And after the first fresh embryo 493 transfer, the post-treatment model provides a revised estimate using treatment related information to 494 personalize the predictions even more. Despite the applicability of the models as counselling tools to 495 inform patients about their prognosis, the McLernon models should not yet be used for decisions on 496 whether or not to withhold fertility treatment. The impact of such model-based decisions on cost-benefit 497 outcomes should be investigated first and proven to be beneficial. To implement the McLernon models as 498 counselling tools in other countries as well, national recalibration is recommended to account for 499 geographical differences in IVF/ICSI treatment.

500 The McLernon models were converted into an online calculator to facilitate the use of the models in 501 clinical practice (https://w3.abdn.ac.uk/clsm/opis). As the original pre-treatment model overestimates 502 cumulative LBR for couples in the Netherlands, conversion of the recalibrated pre-treatment model into a 503 new online calculator is needed for implementation in Dutch clinical practice. This tailored online 504 calculator can then provide accurate and up to date predictions for couples and clinicians in the 505 Netherlands. Ultimately, the online calculator will be offered for implementation on the websites of the 506 Dutch Patient Association for people with fertility problems 'Freya' and the Dutch Association of 507 Obstetrics and Gynaecology (NVOG) to increase the accessibility of the models.

508 *Research implications* 

509 Following this external validation study, future studies could focus on the impact of introducing the 510 McLernon prediction models in clinical practice, and assess changes in patient and clinicians' behaviour 511 and its effects on LBR and cost-effectiveness. In conclusion, after minor recalibration of the pre-treatment model, the McLernon models have proven to be valid in predicting the chance of cumulative live birth after multiple complete treatment cycles in another geographical context and in a more recent time period. Updating the models with AMH, AFC and body weight revealed only a marginal improvement of predictive performance. Following national recalibration, implementation of the McLernon models as counselling tools in clinical practice will provide infertile couples and clinicians with objective and personalized estimates of success over multiple complete IVF/ICSI cycles.

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#### 523 Authors' roles

- T.C.v.T. and S.C.O and all other members from the OPTIMIST study group collected the data. D.J.M.,
  S.B., F.J.M.B. and H.L.T were involved in study conception and study design. J.A.L. and M. J. C. E.
  performed the statistical analysis. J.A.L. drafted the manuscript. J.A.L., M.J.C.E., F.J.M.B. B.W.M.,
  H.L.T interpreted the data. All authors participated to the discussion of the findings and revised the
- 528 manuscript.

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# 539 References

- Akaike H. A new look at the statistical model identification. *IEEE Trans Automat Contr* 1974;19:716–
  723.
- Altman DG, Vergouwe Y, Royston P, Moons KGM. Prognosis and prognostic research: validating a
   prognostic model. *BMJ* 2009;**338**:b605.
- Arvis P, Lehert P, Guivarc'h-Leveque A. Simple adaptations to the Templeton model for IVF outcome
  prediction make it current and clinically useful. *Hum Reprod* 2012;27:2971–2978.
- 546 Berg Brigham K, Cadier B, Chevreul K. The diversity of regulation and public financing of IVF in

Europe and its impact on utilization. *Hum Reprod* 2013;**28**:666–75.

- Boivin J, Bunting L, Collins JA, Nygren KG. International estimates of infertility prevalence and
  treatment-seeking: potential need and demand for infertility medical care. *Hum Reprod*2007;22:1506–12.
- 551 Brandes M, van der Steen JOM, Bokdam SB, Hamilton CJCM, de Bruin JP, Nelen WLDM, Kremer
- JAM. When and why do subfertile couples discontinue their fertility care? A longitudinal cohort
- study in a secondary care subfertility population. *Hum Reprod* 2009;**24**:3127–35.
- Broer SL, van Disseldorp J, Broeze KA, Dolleman M, Opmeer BC, Bossuyt P, Eijkemans MJC, Mol B-

555 WJ, Broekmans FJM, Broer SL, *et al.* Added value of ovarian reserve testing on patient

- characteristics in the prediction of ovarian response and ongoing pregnancy: an individual patient
- data approach. *Hum Reprod Update* 2013;**19**:26–36.
- Cook NR. Statistical Evaluation of Prognostic versus Diagnostic Models: Beyond the ROC Curve. *Clin Chem* 2007;**54**:17–23.
- 560 Coppus SFPJ, van der Veen F, Opmeer BC, Mol BWJ, Bossuyt PMM. Evaluating prediction models in

561	reproductive medicine. Hum Reprod 2009;24:1774–1778.
562	Datta J, Palmer MJ, Tanton C, Gibson LJ, Jones KG, Macdowall W, Glasier A, Sonnenberg P, Field N,
563	Mercer CH, et al. Prevalence of infertility and help seeking among 15 000 women and men. Hum
564	<i>Reprod</i> 2016; <b>31</b> :2108–2118.
565	de Neubourg D, Bogaerts K, Blockeel C, Coetsier T, Delvigne A, Devreker F, Dubois M, Gillain N,
566	Gordts S, Wyns C. How do cumulative live birth rates and cumulative multiple live birth rates over
567	complete courses of assisted reproductive technology treatment per woman compare among
568	registries? <i>Hum Reprod</i> 2016; <b>31</b> :93–99.
569	Dhillon RK, Smith PP, Malhas R, Harb HM, Gallos ID, Dowell K, Fishel S, Deeks JJ, Coomarasamy A.
570	Investigating the effect of ethnicity on IVF outcome. <i>Reprod Biomed Online</i> 2015; <b>31</b> :356–363.
571	Dutch Society of Obstetrics and Gynaecology (NVOG). Landelijke Netwerkrichtlijn Subfertiliteit. 2010.
572	Available at: http://nvog-documenten.nl.
573	Dyer S, Chambers GM, de Mouzon J, Nygren KG, Zegers-Hochschild F, Mansour R, Ishihara O, Banker
574	M, Adamson GD. International Committee for Monitoring Assisted Reproductive Technologies
575	world report: Assisted Reproductive Technology 2008, 2009 and 2010. Hum Reprod 2016;31:1588-
576	1609.
577	Glujovsky D, Farquhar C, Quinteiro Retamar AM, Alvarez Sedo CR, Blake D. Cleavage stage versus
578	blastocyst stage embryo transfer in assisted reproductive technology. Cochrane Database Syst Rev
579	2016:CD002118.
580	Harrell FE. Regression Modeling Strategies: With Applications to Linear Models, Logistic Regression,
581	and Survival Analysis. New York: Springer-Verlag, 2001.
582	Harrell FE, Lee KL, Mark DB. Multivariable prognostic models: issues in developing models, evaluating

assumptions and adequacy, and measuring and reducing errors. <i>Stat Med</i> 1996;15:361–387.
Leushuis E, van der Steeg JW, Steures P, Bossuyt PMM, Eijkemans MJC, van der Veen F, Mol BWJ,
Hompes PGA. Prediction models in reproductive medicine: a critical appraisal†. Hum Reprod
<i>Update</i> 2009; <b>15</b> :537–552.
Liber AC, Warner KE. Has Underreporting of Cigarette Consumption Changed Over Time? Estimates
Derived From US National Health Surveillance Systems Between 1965 and 2015. Am J Epidemiol
2018; <b>187</b> :113–119.
Malizia BA, Hacker MR, Penzias AS. Cumulative Live-Birth Rates after In Vitro Fertilization. N Engl J
<i>Med</i> 2009; <b>360</b> :236–243.
McLernon DJ, Steyerberg EW, te Velde ER, Lee AJ, Bhattacharya S. Predicting the chances of a live
birth after one or more complete cycles of in vitro fertilisation: population based study of linked
cycle data from 113 873 women. <i>BMJ</i> 2016; <b>355</b> :i5735.
Moons KGM, Kengne AP, Grobbee DE, Royston P, Vergouwe Y, Altman DG, Woodward M. Risk
prediction models: II. External validation, model updating, and impact assessment. Heart
2012; <b>98</b> :691–8.
Moons KGM, Kengne AP, Woodward M, Royston P, Vergouwe Y, Altman DG, Grobbee DE. Risk
prediction models: I. Development, internal validation, and assessing the incremental value of a new
(bio)marker. <i>Heart</i> 2012; <b>98</b> :683–90.
National Institute for Health and Care Excellence (NICE). Fertility problems: assessment and treatment.
Clinical guideline. 2013. Available at: https://www.nice.org.uk.
Nelson SM, Lawlor DA. Predicting live birth, preterm delivery, and low birth weight in infants born from
in vitro fertilisation: A prospective study of 144,018 treatment cycles. <i>PLoS Med</i> 2011;8:e1000386.

605	Oakley L, Doyle P, Maconochie N. Lifetime prevalence of infertility and infertility treatment in the UK:
606	results from a population-based survey of reproduction. <i>Hum Reprod</i> 2008;23:447–450.
607	Olivius C, Friden B, Borg G, Bergh C. Why do couples discontinue in vitro fertilization treatment? A
608	cohort study. Fertil Steril 2004;81:258–61.
609	Oudshoorn SC, van Tilborg TC, Eijkemans MJC, Oosterhuis GJE, Friederich J, van Hooff MHA, van
610	Santbrink EJP, Brinkhuis EA, Smeenk JMJ, Kwee J, et al. Individualized versus standard FSH
611	dosing in women starting IVF/ICSI: an RCT. Part 2: The predicted hyper responder. Hum Reprod
612	2017; <b>32</b> :2506–2514.
613	Pandian Z, Marjoribanks J, Ozturk O, Serour G, Bhattacharya S. Number of embryos for transfer
614	following in vitro fertilisation or intra-cytoplasmic sperm injection. Cochrane database Syst Rev
615	2013; <b>7</b> :CD003416.
616	Rajkhowa M, Mcconnell A, Thomas GE. Reasons for discontinuation of IVF treatment: a questionnaire
617	study. <i>Hum Reprod</i> 2006; <b>21</b> :358–363.
618	Rossi B V, Berry KF, Hornstein MD, Cramer DW, Ehrlich S, Missmer SA. Effect of Alcohol
619	Consumption on In Vitro Fertilization. <i>Obstet Gynecol</i> 2011; <b>117</b> :136–142.
620	Rubin DB. Multiple Imputation for Nonresponse in Surveys. In: John Wiley & Sons, 2004.
621	Smeenk JM, Stolwijk AM, Kremer JA, Braat DD. External validation of the templeton model for
622	predicting success after IVF. Hum Reprod 2000;15:1065-8.
623	Smith ADAC, Tilling K, Lawlor DA, Nelson SM. External Validation and Calibration of IVFpredict: A
624	National Prospective Cohort Study of 130,960 In Vitro Fertilisation Cycles. Sun Q-Y (ed). PLoS
625	<i>One</i> 2015; <b>10</b> :e0121357.
626	Sterne JAC, White IR, Carlin JB, Spratt M, Royston P, Kenward MG, Wood AM, Carpenter JR. Multiple

- 627 imputation for missing data in epidemiological and clinical research: potential and pitfalls. *BMJ*628 2009;**338**:b2393.
- 629 Steyerberg EW. *Clinical Prediction Models: A Practical Approach to Development, Validation, and*630 *Updating*. New York, NY: Springer New York, 2009.
- 631 Stockwell T, Zhao J, Greenfield T, Li J, Livingston M, Meng Y. Estimating under- and over-reporting of
- drinking in national surveys of alcohol consumption: identification of consistent biases across four
  English-speaking countries. *Addiction* 2016;**111**:1203–1213.
- 634 Templeton A, Morris JK, Parslow W. Factors that affect outcome of in-vitro fertilisation treatment.
- 635 *Lancet* 1996;**348**:1402–1406.
- te Velde ER, Nieboer D, Lintsen AM, Braat DDM, Eijkemans MJC, Habbema JDF, Vergouwe Y.
- 637 Comparison of two models predicting IVF success; the effect of time trends on model performance.
   638 *Hum Reprod* 2014;29:57–64.
- van Buuren S, Groothuis-Oudshoorn K. MICE : Multivariate Imputation by Chained Equations in R. J *Stat Softw* 2011;45:1–67.
- van Loendersloot L, Repping S, Bossuyt PMM, van der Veen F, van Wely M. Prediction models in in
  vitro fertilization; where are we? A mini review. *J Adv Res* 2014;**5**:295–301.
- van Loendersloot LL, van Wely M, Repping S, van der Veen F, Bossuyt PMM. Templeton prediction
- model underestimates IVF success in an external validation. *Reprod Biomed Online* 2011;22:597–
  602.
- 646 van Tilborg TC, Eijkemans MJ, Laven JS, Koks CA, de Bruin JP, Scheffer GJ, van Golde RJ, Fleischer
- 647 K, Hoek A, Nap AW, *et al.* The OPTIMIST study: optimisation of cost effectiveness through
- 648 individualised FSH stimulation dosages for IVF treatment. A randomised controlled trial. *BMC*

649	Womens	Health	2012;12:29
-----	--------	--------	------------

- van Tilborg TC, Oudshoorn SC, Eijkemans MJC, Mochtar MH, van Golde RJT, Hoek A, Kuchenbecker
  WKH, Fleischer K, de Bruin JP, Groen H, *et al.* Individualized FSH dosing based on ovarian reserve
  testing in women starting IVF/ICSI: a multicentre trial and cost-effectiveness analysis. *Hum Reprod*2017;32:2485–2495.
  van Tilborg TC, Torrance HL, Oudshoorn SC, Eijkemans MJC, Koks CAM, Verhoeve HR, Nap AW,
- 656 starting IVF/ICSI: an RCT. Part 1: The predicted poor responder. *Hum Reprod* 2017;**32**:2496–2505.

Scheffer GJ, Manger AP, Schoot BC, et al. Individualized versus standard FSH dosing in women

- 657 Waylen AL, Metwally M, Jones GL, Wilkinson AJ, Ledger WL. Effects of cigarette smoking upon
- clinical outcomes of assisted reproduction: a meta-analysis. *Hum Reprod Update* 2009;**15**:31–44.
- Wong KM, Mastenbroek S, Repping S. Cryopreservation of human embryos and its contribution to
  in vitro fertilization success rates. *Fertil Steril* 2014;**102**:19–26.
- 661 Zegers-Hochschild F, Adamson GD, Dyer S, Racowsky C, de Mouzon J, Sokol R, Rienzi L, Sunde A,
- 662 Schmidt L, Cooke ID, *et al.* The International Glossary on Infertility and Fertility Care, 2017†‡§.
- 663 *Hum Reprod* 2017;**32**:1786–1801.

664

## 666 Figure legends

Figure 1: Flow chart presenting the numbers (%) of live birth, treatment continuation and discontinuation
over six complete cycles in the OPTIMIST and HFEA databases (McLernon *et al.*, 2016).

669

Figure 2: Calibration plots showing the association between the calculated and observed cumulative live
birth rates over 3 complete IVF/ICSI cycles in the OPTIMIST cohort for a) the *original pre-treatment model* as described by McLernon et al (McLernon *et al.*, 2016) b) *recalibrated pre-treatment model* with
adjustment of the intercept.

674

Figure 3: Calibration plot showing the association between the calculated and observed cumulative live
birth rates over 3 complete IVF/ICSI cycles in the OPTIMIST cohort for the *original post-treatment model* as described by McLernon (McLernon *et al.*, 2016).

678

Figure 4: Example of the *recalibrated pre-treatment model* predicting the cumulative probability of a
live birth up to three complete ICSI cycles for a woman with primary infertility caused by a male factor,
aged 30 or 40 years with an infertility duration of two or five years.

682

**Figure 5:** Example of the with AMH, AFC and body weight *updated pre-treatment model* predicting the cumulative probability of a live birth up to three complete ICSI cycles for a woman with two years of primary infertility caused by a male factor, aged 30 or 40 years, a total body weight of 70 kilograms, with an AMH of 2.0 or 0.5 ng/mL and an AFC of 15 or 7.

688	Figure 6: Example of the <i>post-treatment model</i> predicting the cumulative probability of a live birth up to
689	three complete ICSI cycles for a woman with two years of primary infertility caused by a male factor,
690	aged 30 or 40 years, with 5 or 10 oocytes retrieved, a cleavage stage single embryo transfer, with or
691	without embryo cryopreservation.

693 Supplementary Figure 1. Plots showing the adjusted relation between the predictors included in the
 694 *updated McLernon pre-treatment model* and the probability of a live birth after IVF/ICSI treatment.

695 Predictor; linear predictor (XB) of the original pre-treatment model as described by McLernon
696 (McLernon et al. 2016), Weight; female body weight in kg, AFC; antral follicle count (2-10mm), AMH;
697 anti-Müllerian hormone (ng/mL)

698

699	Supplementary Figure 2. Plots showing the adjusted relation between the predictors in the <i>updated</i>
700	McLernon post-treatment model and the probability of a live birth after IVF/ICSI treatment.

701 Predictor: linear predictor (XB) of the original post-treatment model as described by McLernon

702 (McLernon et al 2016); AFC; antral follicle count (2-10mm), AMH; anti-Müllerian hormone (ng/mL)

703

704



Figure 1: Flow chart presenting the numbers (%) of live birth, treatment continuation and discontinuation over six complete treatment cycles in the OPTIMIST and HFEA databases (McLernon et al., 2016).

209x297mm (300 x 300 DPI)



152x76mm (300 x 300 DPI)



152x152mm (300 x 300 DPI)



Figure 4: Example of the recalibrated pre-treatment model predicting the cumulative probability of live birth up to three complete ICSI cycles for a woman with primary infertility caused by a male factor, aged 30 or 40 years with an infertility duration of two or five years.

152x166mm (300 x 300 DPI)



Figure 5: Example of the with AMH, AFC and body weight updated pre-treatment model predicting the cumulative probability of live birth up to three complete ICSI cycles for a woman with two years of primary infertility caused by a male factor, aged 30 or 40 years, a total body weight of 70 kilograms, with an AMH of 2.0 or 0.5ng/mL and an AFC of 15 or 7.

152x166mm (300 x 300 DPI)



152x166mm (300 x 300 DPI)

# Tables

**Table I** Characteristics of patient and treatment variables included as predictors in the development sample (HFEA cohort) and the validation sample (OPTIMIST cohort) (McLernon *et al.*, 2016). <del>Unless stated otherwise data are n (%).</del>

Characteristics	<b>HFEA cohort</b>	<b>OPTIMIST</b> cohort	Missing
			values in
			OPTIMIST
No of women	112 872	1 511	conort (%)
No of complete cycles	184 269	2 881	
to of complete cycles	101209	2 001	
Patient characteristics			
Age (years), mean (SD)	34.1 (5)	33.5 (5)	2 (0.1)
Duration of infertility (years), median (IQR)	4 (3-6)	2 (2-3)	18 (1.2)
No previous pregnancy in couple <del>,</del>	75 541 (66)	917 (61)	2 (0.1)
Cause of infertility:	26 545 (22)	150 (11)	
- Tubal factor <del>)</del>	26 545 (23)	158 (11)	
- Male factor	49 753 (44)	839 (56)	
- Anovulatory	15 942 (14)	NA by protocol	
- Endometriosis	7 590 (7)	60 (4)	
- Unexplained	32 693 (29)	521 (35)	
Body weight (kg), mean (SD)	NA	69.5 (13)	36 (2.4)
Anti-Müllerian hormone (ng/mL), median (IQR)	NA	1.9 (1-3)	169 (11.2)
Antral follicle count (2-10mm), median (IQR)	NA	13 (9-18)	
Treatment characteristics of first completed			
cvcle			
IVF	67 511 (59)	830 (55)	
ICSI	46 362 (41)	681 (45)	
No of oocytes retrieved, median (IQR)	8 (5-13)	$8(5-13)^{a}$	1 (0.1)
No of embryos created, median (IQR)	5 (2-8)	$4(2-7)^{a}$	4 (0.3)
No of embryos frozen, median (IQR)	0 (0-1)	$1 (0-3)^{a}$	6 (0.5)
Cryopreservation of embryos	28 950 (25)	726 (48)	
Fresh embryo transfer: stage and no. of			24 (1.6)
transferred embryos:	0.249 (9)	1.004 ((())	
- Cleavage stage SET	9 248 (8)	1 004 (66)	
- Cleavage stage DET	75 701 (66)	125 (8)	
- Cleavage stage TET	8 649 (8)	4 (0.3)	
- Blastocyst stage SET	662 (1)	NA	
- Blastocyst stage DET	2 960 (3)	NA	
- Blastocyst stage TET <del>)</del>	130 (0.1)	NA	
- No transfer	15 501 (14)	354 (23)	

<u>Data are presented as number (%) unless Unless stated-otherwise specified. data are n (%).</u> IQR; interquartile range, NA; not available, SET; single embryo transfer, DET; double embryo transfer, TET; triple embryo transfer. <u>a) Median is calculated over 1293 women who had an ovarian follicle aspiration.</u>



Supplementary Figure 1. Plots showing the adjusted relation between the predictors included in the updated McLernon pre-treatment model and the probability of a live birth after IVF/ICSI treatment. # + # + Predictor; linear predictor (XB) of the original pre-treatment model as described by McLernon (McLernon et al. 2016), Weight; female body weight in kg, AFC; antral follicle count (2-10mm), AMH; anti-Müllerian hormone (ng/mL).

152x152mm (300 x 300 DPI)



Supplementary Figure 2. Plots showing the adjusted relation between the predictors in the updated McLernon post-treatment model and the probability of a live birth after IVF/ICSI treatment.!! + !! + Predictor: linear predictor (XB) of the original post-treatment model as described by McLernon (McLernon et al 2016); AFC; antral follicle count (2-10mm), AMH; anti-Müllerian hormone (ng/mL).

152x152mm (300 x 300 DPI)

Supplementary text 1. McLernon pre-treatment model.

**Table showing the predictors in the original McLernon pre-treatment model** (McLernon *et al.*, 2016).

Name predictor	Description	Range of possible values
Age	Female age	18 to 50 years
Duration	How long have you been trying to conceive?	0 to 21 years
Previous	Have you been pregnant before?	1 = No; 0 = Yes
Tubal	Do you have a problem with your tubes?	1 = Yes; 0 = No
Anovulation	Do you have an ovulation problem?	1 = Yes; 0 = No
MaleFactor	Do you have a male factor fertility problem?	1 = Yes; 0 = No
Unexplained	Do you have an unexplained fertility problem?	1 = Yes; 0 = No
Treatment	Which fertility treatment are you planning on having?	1 = ICSI; 0 = IVF

# Original pre-treatment model formulas as described by McLernon et al., 2016).

- 1. For the non-linear relation between Age and the probability of live birth, the following Age1, Age2 and Age3 equations are first calculated and then used in the XB equation below (point 3).
  - Age1 = max((Age-26)/k,0)\*\*3+(11\*max((Age-41)/k,0)\*\*3-(15)\*max((Age-37)/k,0)\*\*3)/4;
  - Age2 =  $max((Age 31)/k, 0)^{*3} + (6^{max}((Age 41)/k, 0)^{*3} (10)^{max}((Age 37)/k, 0)^{*3})/4;$
  - Age3 =  $\max((\text{Age -34})/\text{k},0)**3+(3*\max((\text{Age -41})/\text{k},0)**3-(7)*\max((\text{Age -37})/\text{k},0)**3)/4;$ k=15\*\*(2/3); \*\*means 'to the power of'
- 2. For the non-linear relation between Year and the probability live birth, the following Year1 and Year2 equations are first calculated and then used in the XB equation below (point 3). The value Year= 0 is used for the most up to date predictions.
  - Year1 = max((Year+9)/k,0)\*\*3+((6)\*max((Year)/k,0)\*\*3-(9)\*max((Year+3)/k,0)\*\*3)/(3);
  - Year2 = max((Year+6)/k,0)\*\*3+((3)\*max((Year)/k,0)\*\*3-(6)\*max((Year+3)/k,0)\*\*3)/(3);  $k = 9^{**}(2/3)$ .
- 3. Calculate XB.
  - $$\begin{split} XB = & -0.9948 + 0.0362^{a} + (0.0275*Age) + (-0.1805*Age1) + (0.4553*Age2) + (-1.1990*Age3) \\ & + (-0.0295*Duration) + (-0.0772*Previous) + (-0.0957*Tubal) + (0.0492*Anovulation) + \\ & (-0.1005*MaleFactor) + (0.0602*Unexplained) + (0.2155*Treatment) + (0.0334*Year) + \\ & (-0.0370*Year1) + (0.2173*Year2). \end{split}$$
    - a) To inflate predictions to 2013 an additional 0.0362 is added.
- 4. Calculate the predicted probability of live-birth after the first, second, ...., sixth IVF cycle.

 $\begin{aligned} PCycle1 &= exp(XB)/(1+exp(XB)) \\ PCycle2 &= exp(XB - 0.2394)/(1+exp(XB - 0.2394)) \\ PCycle3 &= exp(XB - 0.4110)/(1+exp(XB - 0.4110)) \\ PCycle4 &= exp(XB - 0.5628)/(1+exp(XB - 0.5628)) \end{aligned}$ 

PCycle5 = exp(XB - 0.7189)/(1+exp(XB - 0.7189))PCycle6 = exp(XB - 0.8138)/(1+exp(XB - 0.8138))

5. Calculate the predicted *cumulative* probability of a live-birth after 1, 2, 3,..., 6 completed IVF cycles:

 $\begin{aligned} & \text{CumPCycle1} = 1-(1-p1) \\ & \text{CumPCycle2} = 1-((1-p1)*(1-p2)) \\ & \text{CumPCycle3} = 1-((1-p1)*(1-p2)*(1-p3)) \\ & \text{CumPCycle4} = 1-((1-p1)*(1-p2)*(1-p3)*(1-p4)) \\ & \text{CumPCycle5} = 1-((1-p1)*(1-p2)*(1-p3)*(1-p4)*(1-p5)) \\ & \text{CumPCycle6} = 1-((1-p1)*(1-p2)*(1-p3)*(1-p4)*(1-p5)*(1-p6)) \end{aligned}$ 

# Supplementary text 2. McLernon post-treatment model.

Name	Description	Range of possible values
predictor		
Age	Female age	18 to 50 years
Duration	How long have you been trying to conceive?	0 to 21 years
Previous	Have you been pregnant before?	1 = No; 0 = Yes
Tubal	Do you have a problem with your tubes?	1 = Yes; 0 = No
Eggs	How many eggs were collected on your first IVF cycle?	(1 to 28)
Treat	Was your first cycle IVF or ICSI?	(1 = ICSI; 0 = IVF)
Cryo	In your first cycle did you have embryos frozen?	(1 = Yes; 0 = No)
Stage	What type of embryo transfer did you have in your first	(No embryos transferred;
C	fresh embryo transfer?	Single cleavage stage;
		Single blastocyst stage;
		Double cleavage stage;
		Double blastocyst stage;
		Triple cleavage stage;
		Triple blastocyst stage)

**Table showing the predictors in the original McLernon post-treatment model** (McLernon *et al.*, 2016).

# Original post-treatment model formulas as described by McLernon et al. (McLernon et al., 2016):

- 1. For the non-linear relation between Age and the probability of live birth, the following Age1, Age2 and Age3 equations are first calculated and then used in the XB equation below (point 4):
  - Age1 = max((Age-26)/k,0)\*\*3+(11\*max((Age-41)/k,0)\*\*3-(15)\*max((Age-37)/k,0)\*\*3)/4;
  - Age2 =  $max((Age 31)/k, 0)^{**3} + (6^{*}max((Age 41)/k, 0)^{**3} (10)^{*}max((Age 37)/k, 0)^{**3})/4;$
  - Age3 = max((Age -34)/k,0)\*\*3+(3\*max((Age -41)/k,0)\*\*3-(7)\*max((Age -37)/k,0)\*\*3)/4; k=15\*\*(2/3), \*\*means 'to the power of'
- 2. For the non-linear relation between Year and the probability of live birth, the following Year1 equation is first calculated and then used in the XB equation below (point 4). The value Year = 0 is used for the most up to date predictions.
  - Year1 = max((Year+8)/k,0)\*\*3+((4)\*max((Year+1)/k,0)\*\*3-(7)\*max((Year+4)/k,0)\*\*3)/(3); k = 7\*\*(2/3).
- 3. For the non-linear relation between Eggs and the probability of live birth, the following Eggs1 equation is first calculated and then used in the XB equation below (point 4):
  - Eggs1=max((Eggs-3)/k,0)\*\*3+((6)\*max((Eggs-18)/k,0)\*\*3-(15)\*max((Eggs-9)/k,0)\*\*3)/(9); k = 15\*\*(2/3).
- 4. Calculate XB

$$\begin{array}{rl} XB = & -1.7564 + 0.0362^{a} + (0.0272*Age) + (-0.1556*Age1) + (0.3812*Age2) + (-1.0184*Age3) \\ & + (-0.0208*Duration) + (-0.0504*Previous) + (-0.2207*Tubal) + (0.0018*Year) + \end{array}$$

(0.0619\*Year1) + (0.0630\*Eggs) + (-0.0479\*Eggs1) + (-0.0968\*Treat) + (0.6490\*Cryo) + Stage<sup>b</sup>

- a) To inflate predictions to 2013 an additional 0.0362 is added.
- b) Stage equals the following values depending on group chosen:
  - If Double cleavage stage then Stage=0;
    - If No embryos transferred then Stage= -1.0842;
    - If Single cleavage stage then Stage= -0.5675;
    - If Single blastocyst stage then Stage= 0.0684;
    - If Double blastocyst stage then Stage= 0.5802;
    - If Triple cleavage stage then Stage= 0.0218;
    - If Triple blastocyst stage then Stage= 0.4547.
- 1. Calculate the predicted probability of live-birth after the first, second, ...., sixth IVF cycle:

 $\begin{aligned} &PCycle1 = exp(XB)/(1+exp(XB)) \\ &PCycle2 = exp(XB - 0.1933)/(1+exp(XB - 0.1933)) \\ &PCycle3 = exp(XB - 0.3537)/(1+exp(XB - 0.3537)) \\ &PCycle4 = exp(XB - 0.5122)/(1+exp(XB - 0.5122)) \\ &PCycle5 = exp(XB - 0.6788)/(1+exp(XB - 0.6788)) \\ &PCycle6 = exp(XB - 0.7666)/(1+exp(XB - 0.7666)) \end{aligned}$ 

2. Calculate the predicted *cumulative* probability of a live-birth after 1, 2, 3,..., 6 complete IVF cycles:

 $\begin{aligned} & \text{CumPCycle1} = 1-(1-p1) \\ & \text{CumPCycle2} = 1-((1-p1)*(1-p2)) \\ & \text{CumPCycle3} = 1-((1-p1)*(1-p2)*(1-p3)) \\ & \text{CumPCycle4} = 1-((1-p1)*(1-p2)*(1-p3)*(1-p4)) \\ & \text{CumPCycle5} = 1-((1-p1)*(1-p2)*(1-p3)*(1-p4)*(1-p5)) \\ & \text{CumPCycle6} = 1-((1-p1)*(1-p2)*(1-p3)*(1-p4)*(1-p5)*(1-p6)) \end{aligned}$ 

Supplementary Text 3. Recalibrated pre-treatment model.

# **Recalibrated pre-treatment model formula**

The included predictors and formulas 1, 2, 4 and 5 are unchanged to the original McLernon pretreatment model (see Supplementary Text 1)

- 3. Calculate XB.
  - $$\begin{split} XB = & -0.3474^{a} 0.9948 + 0.0362^{b} + (0.0275^{*}Age) + (-0.1805^{*}Age1) + (0.4553^{*}Age2) + \\ & (-1.1990^{*}Age3) + (-0.0295^{*}Duration) + (-0.0772^{*}Previous) + (-0.0957^{*}Tubal) + \\ & (0.0492^{*}Anovulation) + (-0.1005^{*}MaleFactor) + (0.0602^{*}Unexplained) + \\ & (0.2155^{*}Treatment) + (0.0334^{*}Year) + (-0.0370^{*}Year1) + (0.2173^{*}Year2). \end{split}$$
    - a) To recalibrate the pre-treatment model, 0.3474 is subtracted from the intercept.
    - b) To inflate predictions to 2013 an additional 0.0362 is added.