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A nested case-control study of adjuvant hormonal therapy persistence and compliance, and early breast cancer recurrence in women with stage I–III breast cancer

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Background: Non-persistence and non-compliance are common in women prescribed hormonal therapy for breast cancer, but little is known about their influence on recurrence.

Methods: A nested case–control study of associations between hormonal therapy non-persistence and non-compliance and the risk of early recurrence in women with stage I–III breast cancer was undertaken. Cases, defined as women with a breast cancer recurrence within 4 years of hormonal therapy initiation, were matched to controls (1:5) by tumour stage and age. Conditional logistic regression was used to examine associations between early recurrence and hormonal therapy non-persistence and non-compliance.

Results: Ninety-four women with breast cancer recurrence were matched to 458 controls. Women who were non-persistent (\geq 180 days without hormonal therapy) had a significantly increased adjusted recurrence odds ratio (OR) of 2.88 (95%Cl 1.11, 7.46) compared with persistent women. There was no significant association between low compliance (OR 1.30; 95% Cl 0.74, 2.30) and breast cancer recurrence.

Conclusion: Hormonal therapy non-persistence is associated with a significantly higher risk of early recurrence in women with stage I–III oestrogen receptor (ER)-positive breast cancer. This finding is consistent with results from randomized studies of hormonal therapy treatment duration and suggests that interventions to target modifiable risk factors for non-persistence are required.

Adjuvant hormonal therapy reduces the annual breast cancer recurrence rate for women with oestrogen receptor (ER)-positive early breast cancer by up to 50%, with at least 5, and up to 10, years of treatment required to achieve the optimal benefit (EBCTG, 2005; Davies *et al*, 2013). Despite the efficacy of hormonal therapy, as many as one in seven women prescribed adjuvant treatment will have a breast cancer recurrence within 5 years of treatment

initiation (EBCTG, 2005), with evidence suggesting a peak in recurrences occurring at 2–3 years post diagnosis (Saphner *et al*, 1996; Debled *et al*, 2007; Kennecke *et al*, 2008; Mansell *et al*, 2009). Various tumour-related factors—including large size, positive lymph node status and high grade—have been identified as predictors of early breast cancer recurrence (Saphner *et al*, 1996; Debled *et al*, 2007; Kennecke *et al*, 2008; Mansell *et al*, 2009).

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Early treatment discontinuation (non-persistence) and sub-optimal treatment execution (non-compliance) are both common in women prescribed hormonal therapies for breast cancer.(Partridge et al, 2003; Barron et al, 2007) Many women cite treatment side effects as the primary reason for not taking their hormonal therapy as prescribed (Grunfeld et al, 2005; Henry et al, 2012). Studies indicate that up to 30% of women will discontinue hormonal treatments within 3-5 years of initiation (Barron et al, 2007; Owusu et al, 2008), and that as many as 20% of women regularly omit at least one in five of their prescribed doses while on treatment (Partridge et al, 2003). A small number of studies have suggested that reduced hormonal therapy exposure, due to either non-persistence or non-compliance, is associated with an increased risk of mortality (McCowan et al, 2008; Dezentjé et al, 2010; Hershman et al, 2011; Weaver et al, 2012; Makubate et al, 2013). However, little is known about the influence of non-persistence and non-compliance on the risk of early breast cancer recurrence (Makubate et al, 2013).

The aim of this study was to examine associations between hormonal therapy non-persistence and non-compliance and the risk of early recurrence in women with ER-positive breast cancer, in analyses adjusted for other predictors of recurrence.

SUBJECTS AND METHODS

Setting and data sources. The study was conducted using patient records from the National Cancer Registry Ireland (NCRI), which are linked to prescription dispensing data from Ireland's Primary care reimbursement services (PCRS) pharmacy claims database. Patient records from the PCRS and NCRI databases were linked using probabilistic matching (AutoMatch, Matchware Technologies, Inc., Silver Spring, MD, USA). NCRI records detailed information on all incident cancers diagnosed in the population usually resident in Ireland. Information on patient characteristics, tumour details, treatment received and death is collected by trained hospital-based tumour registration officers (TRO) from multiple sources including pathology and radiology reports, medical records and death certificates. The accuracy and completeness of NCRI data has been evaluated and described (NCRI, 2012). The PCRS pharmacy claims database is an administrative database of electronic claims from pharmacies for financial reimbursement of dispensed medications. The PCRS is responsible for reimbursement of claims made under the General Medical Services (GMS) community drug scheme, which provides universal healthcare to approximately one-third (1.4 million) of the Irish population. Eligibility for the GMS scheme is through means test and age $(\geq 70 \text{ years})$. The GMS population is therefore older and more socioeconomically deprived than the full Irish population. The PCRS pharmacy claims database records details of all prescription drugs dispensed to patients with eligibility for the scheme. Drugs are coded according to WHO-ATC classifications (WHO, 2010). The use for research of anonymised data held by the NCRI is covered by the Health (Provision of Information) Act 1997.

Study design. A case–control study design, nested within a cohort was used. Women between the ages of 40 and 80 years were eligible for inclusion in the study cohort if they had a diagnosis of stage I–III, ER-positive breast cancer (ICD-10 C50) (Greene *et al*, 2002), between 1 January 2002 and 31 December 2006; had continuous GMS eligibility from at least 1 year prior to diagnosis; had received tumour-directed surgery within 1 year of breast cancer diagnosis (lumpectomy or mastectomy; Appendix 1); and had filled at least one prescription for hormonal therapy within 1 year of breast cancer diagnosis and at least 90 days prior to the end of follow-up. Women were excluded if they had a prior invasive cancer other than non-melanoma skin cancer. Patients were followed up until

breast cancer recurrence, death from any cause, 31 December 2007 or 4 years from hormonal therapy initiation, whichever occurs first.

Covariate definitions. NCRI data was used to classify women by tumour stage (I, IIa, IIb, IIIa and IIIb-c) (Greene et al, 2002); grade (low, intermediate, high and unspecified); ER, PR, HER-2 status (positive, negative and unspecified; Appendix-1); the receipt of radiation therapy in the year post diagnosis (yes, no); and the receipt of chemotherapy prior to hormonal therapy initiation (yes, no). Information on age at diagnosis (years), smoking status (non-smoker, past, current and unspecified) and socioeconomic status was also available. Socioeconomic status was defined according to census-based National Deprivation Index for the area of residence at diagnosis (5-point scale; 1 least deprived, 5 most deprived) (Kelly and Teljeur, 2007). Prescription refill data was used to identify other, potentially confounding, medication use in the year prior to breast cancer diagnosis (exposed, unexposed; Appendix-1): statins (lipophilic, hydrophilic) and aspirin. The number of medication classes (WHO-ATC level 5) dispensed in the year prior to breast cancer diagnosis was used as a measure of comorbidity (Schneeweiss et al, 2001).

Case-control selection. Cases were defined as women with a local or distant breast cancer recurrence within 4 years of hormonal therapy initiation. The date and site of breast cancer recurrence was identified by TROs from two sources; first, from pathology reports of biopsied breast tumour metastases automatically reported by pathology laboratories to the NCRI, and second, from clinical notes and radiology reports. Controls were matched to cases in a ratio of 5:1 by tumour stage and age (5-year calliper) using incidence density sampling without replacement (Robins *et al*, 1986; Bergstralh and Kosanke, 1995). Controls had to be alive and without recurrence at time of a case's recurrence. The date of recurrence for a case was assigned as the index date for each matched control.

Hormonal therapy medication taking behaviours. All prescriptions for selective oestrogen receptor modulators (SERMs; tamoxifen and toremifene) and aromatase inhibitors (AIs; anastrozole, letrozole and exemestane) dispensed to women from the time of breast cancer diagnosis to the end of follow-up were identified from the PCRS database (Appendix 1). The date of dispensing, type of hormonal therapy and number of days' supply on each prescription were abstracted. Using this data a longitudinal daily history of hormonal therapy availability was assembled for each woman by assigning the days' supply from each prescription to sequential days from the date of dispensing (Peterson et al, 2007). In cases where women switched hormonal therapies these were evaluated as a single contiguous treatment episode. These longitudinal daily histories of hormonal therapy availability were used to calculate three measures of medication-taking behaviour: (i) persistence: the last day of hormonal therapy availability prior to the case/control index date was identified, and the cumulative number of consecutive non-persistent days from this date to the index date was used to classify women as persistent (<180 day gap) or non-persistent (>180 day gap); (ii) compliance: the proportion of days covered up to the first of either the case/control index date or the date of non-persistence was identified for each patient (Benner et al, 2002; Barron et al, 2010); (iii) a second measure of compliance was also calculated; for clarity this is referred to as cumulative exposure: the total quantity of adjuvant hormonal therapy taken by a woman is determined by both the duration of treatment (persistence) and the quality of treatment execution during this time (compliance); to reflect this, the cumulative hormonal therapy exposure was defined as the proportion of days covered up to the case/control index date (i.e. including time after non-persistence in women who had

discontinued treatment). Compliance and cumulative exposure were stratified by tertiles for analyses.

Statistical analysis. The frequency and proportion of cases and controls were tabulated by clinical and sociodemographic covariates and by hormonal therapy persistence, compliance and cumulative exposure categories. Means and percentages for controls were weighted by the inverse number of controls matched to each case. Standardized differences (d) were used to assess balance in measured characteristics between matched cases and controls (Austin, 2008). Univariate and multivariate conditional logistic regression was used to estimate odds ratios (OR) with 95% confidence intervals (CI) for associations between breast cancer recurrence in the 4 years after hormonal therapy treatment initiation and hormonal therapy (i) persistence, (ii) compliance and (iii) cumulative exposure (Rothman et al, 2008). These ORs have a rate ratio interpretation, as controls were matched on person time at risk (Pearce, 1993). Covariates were included in multivariate models based on prior knowledge of clinical predictors of early breast cancer recurrence (tumour grade, PR, HER-2, chemotherapy, comorbidity score) (Saphner et al, 1996; Debled et al, 2007; Kennecke et al, 2008; Mansell et al, 2009), demographic predictors of breast cancer outcomes (smoking status, socioeconomic status) (Bradley et al, 2002; Braithwaite et al, 2012) and drug exposures previously associated with breast cancer outcomes (statins, aspirin) (Holmes et al, 2010; Ahern et al, 2011). To avoid over fitting, backwards stepwise elimination of covariates with a threshold α -level of >0.2 was used to select the final multivariate model (Maldonado and Greenland, 1993). All analyses were performed with SAS, version 9.2 (SAS Institute Inc, Cary, NC, USA). Results were considered statistically significant at a two-sided α -level of 0.05.

Sensitivity analyses. Probabilistic sensitivity analyses were undertaken to estimate the potential impact of misclassification of breast cancer recurrence on the primary analysis using the SAS macro SENSMAC available at https://sites.google.com/site/biasanalysis/ sensmac (Lash and Fink, 2003; Fox *et al*, 2005; Lash *et al*, 2009). The parameters for these sensitivity analyses were specified based on (i) estimates from validation studies in external populations using similar methods for ascertainment of recurrence, (ii) knowledge of recurrence assessment practices by NCRI tumour registration officers and (iii) numerical constraints of the data (Jurek *et al*, 2009). The identification of breast cancer recurrence by medical record abstraction has been shown to have high specificity (96%) but imperfect sensitivity (90%) (Lash et al, 2007; Thwin et al, 2007; Bosco et al, 2009). Using these values to guide parameter estimates it was assumed that recurrence assessment by NCRI TROs would have a lower sensitivity but similar specificity. The influence of non-differential misclassification of recurrence was then evaluated (5000 simulations) with sensitivity and specificity drawn from the same trapezoidal distributions. The possible range for these distributions could not be evaluated from prior data and they were conservatively set with a wide dispersion at 0.70-0.90 for sensitivity (mode 0.75-0.85) and 0.94-1.00 for specificity (mode 0.96-0.98). In differential analyses it was assumed a higher sensitivity and specificity would be achieved for women with higher compliance/persistence. A correlation of 0.8 was specified for the similarity between sensitivity and specificity. Finally, the distributions of sensitivity and specificity were adjusted upwards so that no more than 10% of simulations produced impossible corrected data (i.e. negative counts). The final parameters used are presented in Table 3. In addition to these probabilistic sensitivity analyses, analyses were also repeated including patients of any age.

RESULTS

A flow chart outlining the cohort selection is presented in Figure 1. The characteristics of women included in the cohort (n = 1376) are presented in Table 1. The mean age of women at diagnosis was 65.0 years. Radiation and chemotherapy were received by 68.0% and 32.6% of women, respectively. Aromatase inhibitors were prescribed as first-line hormonal therapy to 50.2% of women. During follow-up, 44.3% of patients switched from a SERM to an AI and 8.7% switched from an AI to a SERM. In addition, 6.4% of women were non-persistent with treatment at the time of recurrence (≥ 180 day gap in treatment); the mean hormonal therapy compliance rate while persistent with treatment was 90.0% (s.d. = 15.8%); the mean cumulative exposure rate was 86.8\% (s.d. = 20.8%). A total of 95 women had a breast cancer recurrence within 4 years of hormonal therapy initiation, translating into an overall recurrence rate of 25.6 per 1000 women-years. The most frequent recurrent metastatic site was bone (48.4%), followed by liver (11.6%), lung (8.4%), brain (7.4%) and local recurrence (7.4%).

In the case–control analysis, 94 of the women with recurrence were successfully matched to 458 controls (89 cases had 5 controls each, 1 case had 4 controls, 2 cases had 3 controls, 1 case had 2



Figure 1. Flow chart showing study cohort inclusion and exclusion criteria.

Table 1. Characteristics of full cohort, breast cancer recurrence cases and matched controls					
Characteristic	Full cohort (n = 1376)	Cases ^a (n = 94)	Controls ^{a,b} (<i>n</i> = 458)		
Patient details					
Age—mean (s.d.)					
Years	65.0 (10.2)	64.3 (10.0)	64.3 (10.1)		
Comorbidity—mean (s.d.)		1			
Drug classes	8.4 (6.3)	9.0 (7.1)	8.1 (6.1) ^c		
Smoking status (%)					
Current	319 (23.2)	23 (24.5)	98 (21.6)		
Never	706 (51.3)	40 (42.6)	241 (52.6)°		
Unspecified	191 (13.9)	16 (17.0)	66 (14.4)		
Deprivation index (%)		1			
1—least deprived	183 (13.3)	14 (14.9)	57 (12.1)		
2	139 (10.1)	11 (11.7)	49 (10.5)		
3	168 (12.2)	10 (10.6)	62 (13.2)		
4 5—most deprived	230 (17.3) 542 (39.4)	30 (31 9)	183 (41 3) ^c		
Not assigned	106 (7.7)	10 (10.6)	33 (7.0) ^c		
Tumour details					
Tumour stage (%) ^d		_			
1	446 (32.4)	13 (13.8)	65 (13.8)		
lla / llb	447/316 (32.5/23.0)	28/29 (29.8/30.9)	140/145 (29.8/30.9)		
	79/00 (5.7/0.4)	11/15 (11.7/15.0)	467.60 (11.7715.6)		
Tumour grade (%)					
Low	173 (12.6)	6 (6.4)	42 (9.0)		
Intermediate	743 (54.0)	41 (43.6)	246 (53.2)°		
High Unspecified	342 (24.9) 118 (8.6)	38 (40.4)	45 (9.6)		
	110 (0.0)	7 (7.0)	43 (7.0)		
PR (%)		1			
Negative	197 (14.3)	20 (21.3)	68 (14.8)		
Positive	919 (66.8) 260 (18 9)	54 (57.4) 20 (21.3)	300 (65.6)*		
HER-2 (%)	200 (10.7)	20 (21.3)	10 (17.7)		
Negative	707 (57 0)	50 (52 2)	258 (54.7)		
Positive	128 (9.3)	11 (11.7)	43 (9.1)		
Unspecified	451 (32.8)	33 (35.1)	157 (34.2)		
Treatment details					
Chemotherapy (%) ^e	449 (32.6)	39 (41.7)	187 (41.5)		
Radiation (%)	936 (68.0)	60 (63.8)	318 (69.6)		
Tamavifan		50 ((2.0)	0/4/5/ 0)		
i amoxiten Toremifene	685 (49.8) 0 (0 0)	57 (62.8)	261 (56.8)		
Anastrozole	586 (42.6)	29 (30.9)	166 (36.6)		
Letrozole	99 (7.2)	6 (6.4)	28 (6.0)		
Exemestane	6 (0.4)	0 (0.0)	3 (0.6)		
Other drug exposures (%)					
Aspirin	357 (25.9)	23 (24.5)	117 (25.7)		
Statin hydrophilic Statin lipophilic	302 (21.9)	18 (19.1)	89 (19.2) 17 /2 4) ^c		
statin ilpoprinic	30 (2.0)	1 (1.1)	17 (3.0)		

Hormonal therapy persistence and compliance

Table 1. (Continued)					
Characteristic	Full cohort (<i>n</i> = 1376)	Cases ^a (n = 94)	Controls ^{a,b} (n = 458)		
Recurrence details					
Follow-up—mean (s.d.)					
Years	2.70 (1.14)	1.86 (1.01)	1.86 (1.04)		
Breast cancer recurrences (%)	95 (6.9)	_	_		
Woman-years	3716				
Breast cancer recurrence rate					
Recurrence/1000 years	25.6				
Recurrence sites (%) ⁹					
Liver	11 (11.6)	_	_		
Lung	8 (8.4)	—	—		
Bone	46 (48.4)	—	—		
Brain	7 (7.4)	—	—		
Other sites	20 (21.1)	—	—		
Unspecified	11 (11.6)	—	_		

Abbreviations: HER-2=human epidermal growth factor receptor 2; PR=progesterone receptor; s.d.=standard deviation.

^aCases: women with a breast cancer recurrence within 4 years of hormonal therapy initiation. Controls: women without a breast cancer recurrence at the time of a matched case's recurrence. Controls were randomly matched to cases in a ratio of 5:1, on tumour stage at diagnosis and age within a calliper of 5 years, using incidence density sampling without replacement. ^bMeans and percentages for controls are weighted by the inverse number of controls matched to each case; 89 cases had 5 matched controls, 1 case had 4 matched controls, 2 cases had 3

matched controls, 1 case had 2 matched controls and 1 case had 1 matched control.

^cWeighted standardized difference (d) between cases and controls >0.1.

^dAJCC Cancer Staging Manual 6th Edition. Springer (Greene *et al*, 2002).

^eChemotherapy received prior to hormonal therapy initiation.

^fRadiation received in the year post diagnosis.

controls, 1 case had 1 control and 1 case had 0 controls). The characteristics of these cases and matched controls are presented in Table 1. Cases were more likely to have tumour characteristics previously associated with breast cancer recurrence such as higher grade or negative progesterone receptor status. In multivariate analyses of hormonal therapy compliance and persistence, adjusted for prognostic tumour and patient characteristics, breast cancer recurrence ORs were increased for women in the nonpersistent group and the lowest compliance tertile, although only associations with non-persistence reached formal statistical significance. Women who were non-persistent with treatment had a significantly increased adjusted recurrence OR of 2.88 (95% CI 1.11, 7.46; 8 cases, 14 controls; Table 2) in comparison to women who persisted with treatment. Women in the lowest compliance tertile had a non-significant adjusted breast cancer recurrence OR of 1.30 (95% CI 0.74, 2.30; 36 cases, 148 controls), in comparison to women with highest compliance (Table 2). In analyses of total cumulative hormonal therapy exposure, from treatment initiation to case/ control index date, the recurrence OR for women with low overall exposure was 1.62 (95% CI 0.91, 2.88; 43 cases, 161 controls; Table 2); this was not formally significant (P = 0.11).

Sensitivity analyses. The results from probabilistic sensitivity analyses correcting for possible non-differential and differential misclassification of breast cancer recurrence are presented in Table 3. In these sensitivity analyses the median OR estimates for low persistence, compliance and cumulative exposure, based on 5000 simulations, were similar to estimates obtained from the primary analyses. The 95% OR simulation limits for associations between low persistence and breast cancer recurrence did not include the null, which is compatible with results from the primary analysis. Additionally, in a analyses including patients of any age the adjusted recurrence ORs were not considerably different from those from the main analysis: non-persistence OR = 2.46 (95% CI

1.00, 6.07; 8 cases, 17 controls); lowest compliance tertile OR = 1.60 (95% CI 0.92, 2.77; 41 cases, 164 controls); lowest overall exposure tertile OR = 1.66 (95% CI 0.97, 2.84; 45 cases, 168 controls).

DISCUSSION

In this study of 1376 women with stage I–III ER-positive breast cancer, early breast cancer recurrences were significantly more likely to occur in those who had stopped taking hormonal therapy for \geq 180 days than those who persisted with treatment. In addition, low compliance with treatment appeared to be associated with a moderately increased risk of recurrence, although associations did not reach statistical significance. These results provide evidence, in a 'real-world' clinical practice setting, to support the importance of persistence (and potentially, compliance) with hormonal therapy in the adjuvant treatment of early breast cancer. They are also consistent with the results from randomised controlled trials, which have conclusively demonstrated the benefits of longer-duration adjuvant hormonal therapy for women with early-stage ER-positive breast cancer (EBCTG, 2005; Davies *et al*, 2013).

A small number of prior observational studies have examined associations between hormonal therapy non-persistence and breast cancer recurrence or mortality (Hershman *et al*, 2011; Weaver *et al*, 2012). These have reported mixed results. In a study of 857 women with stage I–III breast cancer, no association was observed between hormonal therapy non-persistence and disease recurrence (OR 1.18, 95% CI 0.76, 1.82) (Weaver *et al*, 2012). The absence of an association may have been due to the definition of non-persistence used; this was assessed within in the first year of treatment only, after which patients could recommence therapy. Conversely, in a larger study of 8769 women with stage I–III breast

Table 2. Numbers and percentages of cases and controls and univariate and multivariate odds ratios for breast cancer recurrence

				Multivariate	
	Cases (%) ^a	Controls (%) ^a	Univariate odds ratio	odds ratio	
Medication-taking behaviour	(n = 94)	(n=458)	(95% CI)	(95% CI) ²	
Persistence and compliance					
Hormonal therapy persistence ^{c,d}					
Persistent	86 (91.5)	444 (96.9)	Ref	Ref	
Non-persistent	8 (8.5)	14 (3.1)	3.00 (1.18, 7.60)	2.88 (1.11, 7.46)	
(>180 day gap)					
Hormonal therapy compliance ^{e,f}					
High (98–100%)	30 (31.9)	154 (33.6)	Ref	Ref	
Intermediate (90–98%)	28 (29.8)	156 (34.1)	0.96 (0.53, 1.71)	0.95 (0.53, 1.71)	
Low (0–90%)	36 (38.3)	148 (32.3)	1.24 (0.71, 2.14)	1.30 (0.74, 2.30)	
Cumulative exposure					
Cumulative hormonal therapy exposure ⁹					
High (98–100%)	24 (25.5)	144 (31.4)	Ref	Ref	
Intermediate (90–98%)	27 (28.7)	153 (33.4)	1.04 (0.57, 1.91)	1.02 (0.55, 1.90)	
Low (0–90%)	43 (45.7)	161 (35.2)	1.60 (0.91, 2.81)	1.62 (0.91, 2.88)	

Abbreviations: CI = confidence interval; Ref: referent group. *P-trend <0.05.

^aCases: women with a breast cancer recurrence within 4 years of hormonal therapy initiation. Controls: women without a breast cancer recurrence at the time of a matched case's recurrence. Controls were randomly matched to cases in a ratio of 5:1, on tumour stage at diagnosis and age within a calliper of 5 years, using incidence density sampling without replacement.

Adjusted for tumour grade (low, intermediate, high and unspecified), progesterone receptor status (positive, negative and unspecified) and comorbidity score.

^cThe number of consecutive non-persistent days from the last day of hormonal therapy availability to the index date, stratified as persistent (<180 day gap) and non-persistent (≥180 day gap). ^dAdjusted for hormonal therapy compliance.

^eThe proportion of days covered up to the first of either the date of non-persistence or the case/control index date, stratified by tertiles.

^fAdjusted for hormonal therapy persistence.

 ${\boldsymbol{\mathsf{g}}}_{\mathsf{The}}$ proportion of days covered up to the case/control index date.

cancer, hormonal therapy non-persistence, defined as a treatment gap of 180 days, was associated with a significantly increased risk of all-cause mortality (HR 1.26, 95% CI 1.09–1.46) (Hershman *et al*, 2011). However, the time-varying nature of treatment persistence does not appear to have been accounted for in these analyses and this result may underestimate the true association between hormonal therapy non-persistence and mortality (Suissa, 2007).

The moderate-albeit non-significant-association between low compliance and increased risk of breast cancer recurrence observed in this study is comparable with the results from a number of prior studies (Dezentjé et al, 2010; Weaver et al, 2012; Makubate et al, 2013). By convention, a compliance threshold of 80% and treatment gap of between 90-180 days have been used to define adjuvant hormonal therapy non-compliance and non-persistence (McCowan et al, 2008; Dezentjé et al, 2010; Hershman et al, 2011; Weaver et al, 2012; Makubate et al, 2013). However, the exact thresholds that define clinically relevant non-compliance and nonpersistence are currently unknown. The results presented here suggest that hormonal therapy treatment discontinuation is associated with a greater risk of breast cancer recurrence than low compliance. Confirmation of these findings and further elucidation of the role that different patterns of missed hormonal therapy doses have in treatment failure will require detailed data, such as that provided by electronic medication event monitoring.

In terms of clinical implications, the results from this study suggest the possibility that interventions aimed at increasing persistence with hormonal therapy could reduce recurrence rates in women with early breast cancer. To date, however, simple educational interventions to improve hormonal therapy compliance and persistence have had limited success (Hadji *et al*, 2012).

er, the exact limitations. The number of cases/controls that contributed to the analysis of persistence was small and the results from this analysis should be interpreted with caution. The assessment of medication-taking behaviour was based upon prescription refill data and may therefore overestimate compliance and persistence in women who fill a prescription but do not take the medication they receive. It was also not possible to validate the data on breast cancer recurrences, or to identify the number of recurrences that were

fill a prescription but do not take the medication they receive. It was also not possible to validate the data on breast cancer recurrences, or to identify the number of recurrences that were biopsy proven, and the possibility that there may have been underascertainment of breast cancer recurrences must be considered. This is unlikely to have greatly influenced the observed associations for the following reasons. First, the case–control methodology employed is appropriate for use in analyses where case ascertainment may be imperfect, in particular where sensitivity may be low but specificity is likely to be very high (Brenner and Savitz, 1990),

It is likely that the systematic development of a theoretically based

intervention to target modifiable risk factors for non-persistence

may be required. The reasons for hormonal therapy non-

persistence were not available for this study. Previous studies have

however reported that side effects strongly influence a woman's

decision to discontinue treatment (Grunfeld *et al*, 2005; Henry *et al*, 2012), suggesting that an intervention might focus on (i) the

early identification of women experiencing side effects; (ii) the

availability of effective supportive pharmacologic and psychologi-

cal care and (iii) the timely switching to alternative hormonal

prospectively collected, population-based, prescription and cancer

registry data. Cases and controls were also sampled from the same

base population and controls were representative of the population

that gave rise to the cases. However, there are a number of

The strengths of this study include the use of high-quality

therapies where appropriate (Henry et al, 2012).

Table 3. Probabilistic sensitivity analyses correcting for non-differential and differential misclassification of breast cancer recurrence						
Sensitivity analysis ^a	Odds ratio 2.5 percentile	Odds ratio median estimate	Odds ratio 97.5 percentile	Change in simulation interval width (<i>b/a</i>) compared with conventional (%)	Proportion of simulations with odds ratio < 1	
Persistence—low (≥180 day gap)						
Non-differential sensitivity analysis ^b	,					
Sensitivity analysis only Sensitivity analysis with random error	2.45 1.39	3.98 4.00	7.78 12.30	- 51.5 35.4	0.010 0.005	
Differential sensitivity analysis ^c						
Sensitivity analysis only Sensitivity analysis with random error	2.68 1.51	4.26 4.29	7.86 12.79	- 53.7 34.1	0.009 0.003	
Compliance–low (0%–90%)						
Non-differential sensitivity analysis ^b						
Sensitivity analysis only Sensitivity analysis with random error	1.00 0.73	1.36 1.35	1.90 2.60	- 37.2 17.5	0.222 0.171	
Differential sensitivity analysis ^c						
Sensitivity analysis only Sensitivity analysis with random error	1.08 0.77	1.46 1.46	2.12 2.81	- 38.5 19.8	0.234 0.124	
Cumulative exposure—low (0%–90%)						
Non-differential sensitivity analysis ^b						
Sensitivity analysis only Sensitivity analysis with random error	1.32 0.92	1.80 1.81	2.61 3.56	– 35.7 25.3	0.048 0.042	
Differential sensitivity analysis ^c						
Sensitivity analysis only Sensitivity analysis with random error	1.43 1.01	1.94 1.96	2.72 3.67	- 40.8 13.1	0.057 0.024	

^a5000 simulations

bNon-differential misclassification of breast cancer recurrence trapezoidal distributions: sensitivity (min=0.70, mode—1=0.75, mode—2=0.85, max=0.90); specificity (min=0.94, mode—1=0.96, mode—2=0.98, max=1.00).

^cDifferential misclassification of breast cancer recurrence trapezoidal distributions: sensitivity: low compliance/persistence/exposure (min = 0.70, mode—1 = 0.75, mode—2 = 0.85, max = 0.90); sensitivity: high compliance/persistence/exposure (min = 0.80, mode—1 = 0.85, mode—2 = 0.95, max = 1.00); specificity: low compliance/persistence/exposure (min = 0.94, mode—1 = 0.96, mode—2 = 0.98, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.97, mode—2 = 0.98, max = 1.00); specificity: low compliance/persistence/exposure (min = 0.85, mode—2 = 0.95, mode—2 = 0.99, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.97, mode—2 = 0.98, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.97, mode—2 = 0.98, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.85, mode—2 = 0.95, mode—2 = 0.99, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.85, mode—2 = 0.95, mode—2 = 0.99, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.95, mode—1 = 0.97, mode—2 = 0.99, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.95, mode—1 = 0.97, mode—2 = 0.99, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.95, mode—1 = 0.97, mode—2 = 0.99, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.95, mode—1 = 0.97, mode—2 = 0.99, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.95, mode—1 = 0.97, mode—2 = 0.99, max = 1.00); specificity: high compliance/persistence/exposure (min = 0.95, mode—1 = 0.97, mode—1 = 0.96, mode—1 = 0.97, mode—1 = 0.97, mode—1 = 0.97, mode—1 = 0.96, mode—1 =

as in this study. Second, the recurrence rate observed in this study (25.6 recurrences/1000 woman-years) is broadly consistent with the recurrence rate observed for women receiving hormonal therapy for ER-positive early breast cancer in a meta-analysis of clinical studies (29.5 recurrences/1000 woman-years) (EBCTG, 2005). Third, probabilistic sensitivity analyses were undertaken to quantify the range of plausible associations in the presence of both differential and non-differential misclassification of recurrences. These indicate that the observed results are unlikely to be influenced greatly by recurrence misclassification within the range of sensitivities and specificities examined. The validity of these sensitivity analyses is however dependent on the ability to accurately specify distributions for the bias parameters, for which there was minimal prior data. One source of residual confounding should be considered as a potential explanation for the observed results. High compliance/persistence behaviour is strongly associated with other healthy behaviours (Curtis et al, 2011); consequently, observed benefits of treatment persistence may be partly attributable to other healthy behaviours (Bonadonna and Valagussa, 1981). Finally, women in this study received a range of different hormonal treatments, each with unique pharmacokinetic/dynamic properties. There was not sufficient statistical power to be able to examine these separately and care should therefore be taken in generalising the results to individual hormonal therapies.

In conclusion, this study found that hormonal therapy nonpersistence was associated with significantly higher risk of breast cancer recurrence in women with stage I–III ER-positive early breast cancer. The results of this analysis are consistent with data from randomized studies of hormonal therapy treatment duration (EBCTG, 2005; Davies *et al*, 2013) and have implications for the development of interventions aimed at increasing persistence with hormonal therapy.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

DISCLAIMER

The Health Research Board Ireland had no role in the study design; in the collection, analysis and interpretation of data; in the writing of the report; or in the decision to submit the paper for publication. The interpretation and reporting of these data are the responsibility of the authors and in no way should be seen as the official policy or interpretation of the National Cancer Registry Ireland or the Irish Health Services Executive Primary Care Reimbursements Services.

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