**Breast Cancer Research** 





# A comparison of five methods of measuring mammographic density: a casecontrol study

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# Abstract

**Background:** High mammographic density is associated with both risk of cancers being missed at mammography, and increased risk of developing breast cancer. Stratification of breast cancer prevention and screening requires mammographic density measures predictive of cancer. This study compares five mammographic density measures to determine the association with subsequent diagnosis of breast cancer and the presence of breast cancer at screening.

**Methods:** Women participating in the "Predicting Risk Of Cancer At Screening" (PROCAS) study, a study of cancer risk, completed questionnaires to provide personal information to enable computation of the Tyrer-Cuzick risk score. Mammographic density was assessed by visual analogue scale (VAS), thresholding (Cumulus) and fully-automated methods (Densitas, Quantra, Volpara) in contralateral breasts of 366 women with unilateral breast cancer (cases) detected at screening on entry to the study (Cumulus 311/366) and in 338 women with cancer detected subsequently. Three controls per case were matched using age, body mass index category, hormone replacement therapy use and menopausal status. Odds ratios (OR) between the highest and lowest quintile, based on the density distribution in controls, for each density measure were estimated by conditional logistic regression, adjusting for classic risk factors.

**Results:** The strongest predictor of screen-detected cancer at study entry was VAS, OR 4.37 (95% CI 2.72–7.03) in the highest vs lowest quintile of percent density after adjustment for classical risk factors. Volpara, Densitas and Cumulus gave ORs for the highest vs lowest quintile of 2.42 (95% CI 1.56–3.78), 2.17 (95% CI 1.41–3.33) and 2.12 (95% CI 1.30–3.45), respectively. Quantra was not significantly associated with breast cancer (OR 1.02, 95% CI 0.67–1.54). Similar results were found for subsequent cancers, with ORs of 4.48 (95% CI 2.79–7.18), 2.87 (95% CI 1.77–4.64) and 2.34 (95% CI 1.50–3.68) in highest vs lowest quintiles of VAS, Volpara and Densitas, respectively. Quantra gave an OR in the highest vs lowest quintile of 1.32 (95% CI 0.85–2.05).

**Conclusions:** Visual density assessment demonstrated a strong relationship with cancer, despite known inter-observer variability; however, it is impractical for population-based screening. Percentage density measured by Volpara and Densitas also had a strong association with breast cancer risk, amongst the automated measures evaluated, providing practical automated methods for risk stratification.

Keywords: Breast density, Case-control, Risk, Cancer, PROCAS

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# Background

High mammographic density, the relative proportion of fibroglandular to fatty tissue in the breast, reduces the effectiveness of mammographic screening [1–4] and increases risk of developing breast cancer [5, 6]. The relationship of density with risk was established using expert visual assessment of film mammograms [7], with computer-assisted methods providing more reproducible estimates [8, 9]. With increasing uptake of full-field digital mammography (FFDM), the association between automated density assessment methods and cancer risk is under investigation [10–12].

The most widely used method of assessing mammographic density in the USA is the Breast Imaging Reporting And Data System (BI-RADS) categorisation, where experts assign mammograms to one of four classes, the upper two being considered "dense" [13]. The class descriptors were changed in 2013 to better identify women whose cancers may be masked by dense parenchymal tissue [14]. Visual assessment of percentage density may be recorded on visual analogue scales (VAS), providing a continuous measure. This yielded a strong relationship with breast cancer risk for film mammograms, with an odds ratio (OR) of approximately 7 for 76–100% density relative to 0-25% [15]. Likewise, Boyd's 6-class categorisation of percent visual density gave a relative risk in the highest category (>75% dense) compared with the lowest of 6.05 (95% confidence interval (CI) = 2.82-12.97) in a case-control study with 354 cases [8]. Cumulus, a semi-automated thresholding method, was developed to improve reproducibility [8, 9] and has a wellestablished relationship with cancer risk [8, 12]. However, this method also requires trained observers, and whilst separating the breast from the mammogram background is reproducible, judgement of the best threshold to separate dense tissue from fat is less so. Boyd category, VAS and Cumulus are all relative, area-based methods, so density estimates can vary depending on breast positioning and patient weight [8, 16]. Weight change disproportionately alters the fatty component of the breast [17] and percentage density measures should be adjusted to take body mass index (BMI) into account [18].

Now that digital mammography is standard throughout the UK, volumetric measures of mammographic density, made by calibrating pixel values in the raw ("for processing") FFDM image using a model of x-ray physics and imaging parameters [19], are now available. These can be expressed either in percentage terms (volumetric percent dense) or as absolute measures of dense and non-dense tissue.

The availability of fully automated density assessment paves the way for risk stratification in screening [20], allowing selection of the most appropriate imaging modality and screening frequency for the individual [21, 22]. The addition of mammographic density to breast cancer risk models based on other risk factors has demonstrated increased predictive power, depending on the method used for density estimation [23–25]. It is therefore important to determine which density methods are suitable for risk-adapted screening; more accurate risk prediction will enable better targeting of risk-reducing interventions including chemoprevention and lifestyle modification [26, 27].

A previous case-control analysis, carried out in London, compared density measured in the unaffected (contralateral) breast in 414 women diagnosed with unilateral breast cancer at one hospital with that of 685 unmatched controls attending routine breast screening. Comparing the highest percentage density quintile with the lowest, and adjusting for age, BMI and reproductive variables, the strongest association with risk of developing breast cancer was for Volpara, with an OR of 8.26 (95% CI 4.28-15.96), followed by Quantra, OR 3.94 (2.26-6.86) and Cumulus, OR 3.38 (2.00-5.72) [13]. However, mammographic density was assessed at the time of detection of cancer, so the ability of density to predict women who would later develop the disease was not assessed. Here we address this by evaluating the association between five mammographic density methods and the presence of cancer at the time of screening, and the association between four mammographic density methods and cancer detected subsequently, either between screening rounds or at a later screen, using data from the Predicting Risk of Cancer At Screening study (PROCAS) [20].

## Methods

## Study design

Women invited to the Greater Manchester Breast Screening Service for routine 3-yearly mammographic screening between October 2009 and March 2015 were also invited to participate in the "Predicting Risk Of Cancer At Screening" (PROCAS) study, which aimed to provide women with a personalised risk estimate of their breast cancer risk based on mammographic density and classic breast cancer risk factors obtained via a questionnaire and quantified by the Tyrer-Cuzick risk score [28]. After October 2012 only women attending their first (prevalent round) screen were invited. At the time of recruitment informed consent was obtained from all participants.

In order to assess density using fully automated methods, the raw FFDM (for processing) image data from GE Senographe Essential mammography systems was obtained. Cancers (invasive and ductal carcinoma in situ) were identified through hospital records or through the North West Cancer Intelligence Service; women who moved out the area were considered ineligible. Two case-control datasets were created. In study 1, cases were women with breast cancer detected at the screen on entry to PROCAS and in study 2 cases were women who were breast cancer free at the screen on entry to PROCAS but had breast cancer detected subsequently, either between screening rounds or at a later screen. In these women we analysed the density of the screen on entry to PROCAS.

Three controls without cancer were matched to each cancer case based on age ( $\pm 12$  months), BMI category (missing, < 24.9, 25.0–29.9, 30+ kg/m<sup>2</sup>), hormone replacement therapy (HRT) use (current vs never/ever) and menopausal status (premenopausal, perimenopausal or postmenopausal). In both studies all controls had a subsequent cancer-free screening mammogram so it was unlikely that early signs of cancer were visible, and in study 2, controls were also matched on year of mammogram at entry.

# Mammographic density measurement Visual estimation of percentage density

Processed FFDM images were displayed on Planar Dome E5 5MP self-calibrating high-resolution monitors. Two of nineteen readers (usually a consultant radiologist or breast physician and an advanced practitioner radiographer) independently recorded density estimates on a paper form showing four 10-cm horizontal VAS, one for each view, labelled 0% and 100% at the ends of the scale. Forms were read using custom software and visual percentage density calculated. VAS readings were averaged between readers and views, and analysed in quintiles and as Boyd categories (0%, >0-10%, >10-25%, >25-50%, > 50–75% and > 75%) [8]. Due to the small number of cases in the highest category (three in study 1 and six in study 2), the top two Boyd categories were combined for analysis. Intra-observer and inter-observer agreement for 120 mammograms randomly selected across deciles of VAS density scores, from the PROCAS study, were assessed by 11 readers, on two occasions, 3 years apart. The majority of readers had excellent intra-observer agreement (intraclass correlation coefficient (ICC) > 0.80), and inter-observer agreement for consistency was excellent (ICC = 0.82) and was substantial for absolute agreement (ICC = 0.69) [29].

## Cumulus

Cumulus (Sunnybrook Health Sciences Centre, Toronto, ON, Canada) density assessment was undertaken by a single reader (JS) trained in August 2010 and validated by a member of the PROCAS team (JW) who had herself been trained by the group that developed the software. Reader performance was validated on test sets of data developed for this purpose by the trainers. Processed FFDM images were analysed. Cumulus was undertaken on a single contralateral mediolateral oblique (MLO) view of a subset of the study 1 dataset comprising 311 screen-detected cancers and their matched controls. The reader was blind to case-control status.

## Quantra™

Quantra version 2.0 (Hologic Inc, Bedford, MA, USA) was used to assess density from the raw FFDM images for each view, each breast and each woman, giving breast and fibroglandular tissue volume (cm<sup>3</sup>), and the dense tissue area as a percentage of breast volume. It also provides a quantized BI-RADS-like score for each view and per breast.

# Volpara™

Volpara Density Algorithm 1.5.0 (Volpara Health Technologies, Wellington, New Zealand) was also used to assess density from the raw FFDM images for each view, giving breast volume and fibroglandular tissue volume (cm<sup>3</sup>) and percentage density by volume. Volpara provided a macro, which produced per-patient results including Volpara Density Grade (VDG 4<sup>th</sup> and 5<sup>th</sup> Edition), designed to correlate with BI-RADS 4<sup>th</sup> and 5<sup>th</sup> Edition [15]. This also computes the percentage density of the two breasts following outlier removal.

## Densitas™

Densitas version 2.0.0 (Densitas Inc, Halifax, NS, Canada) analyses processed FFDM images, giving breast and fibroglandular area (cm<sup>2</sup>) and percentage density by area for each image and per patient. It also produces per-patient measures of BIRADS 4<sup>th</sup> and 5<sup>th</sup> edition [15].

## Statistical methods

In study 1, mammographic density was assessed in the contralateral breast in women with cancer and the breast on the same side in matched controls, whereas in study 2, density was assessed in both breasts at entry to PRO-CAS and the average was used.

Categorical data were compared using the chi-square test for proportions. For ordinal variables, a chi-square test for trend was also conducted. Continuous variables were assessed by the median and Mann-Whitney U test.

The relationship between density assessment and casecontrol status was analysed using conditional logistic regression. Density measures were modelled as quintiles based on the density distributions of controls, and also as continuous measures, transformed to approximately follow a normal distribution (square root transformation for VAS and Cumulus, and a logarithm transformation for Volpara, Quantra and Densitas). Univariate models were fitted initially, and multivariate models fitted to adjust for the logarithm 10-year Tyrer-Cuzick (v.6) risk score. In study 2 we also adjusted for parity, due to imbalance between cases and controls. We also performed an analysis in a subset of women who had been assessed using all density methods to determine which model performed best and differences between models were compared using the likelihood-ratio chi square. The matched concordance (mC) index, a modification of the concordance index (or area under the receiving operator characteristic curve (AUC)) for matched case-control studies, gives an average concordance index within matched groups (where 1.0 would indicate perfect discrimination after allowing for matching factors) with empirical bootstrap confidence intervals [30], was calculated to compare the discrimination performance of risk factors. All p values were two-sided. Analysis was performed in SPSS version 22 [31] and R 3.3.1 [32].

## Results

Of the 57,905 women recruited to PROCAS, raw FFDM image data were available for 44,658 women (77%). Unavailability of raw FFDM images was predominantly due to the use of film mammography initially. There were 1004 cases of cancer occurring after consent up to November 2015, of which 704 were included in the analysis. The excluded women comprised 39 women with a preexisting diagnosis of breast cancer, 13 with synchronous bilateral breast cancer, 118 with film mammograms and 130 with FFDM but for which raw image data was unavailable. Of the 704 women eligible for the analysis, 366 were women with breast cancer detected at the screen on entry to PROCAS (study 1) and 338 were women who were found to be breast cancer free at the screen on entry to PROCAS but had breast cancer detected subsequently, either between screening rounds or at a later screen (study 2). Of the latter, 114 women developed an interval cancer within 5–46 months of entry (IQR 13–31) and 224 women had breast cancer detected at a subsequent screen 17-55 months after entry (IQR 35-38).

Matching was satisfactory for both studies (Table 1). There was a difference in 10-year Tyrer-Cuzick score, with the score higher in cases (study 1, 2.95 vs 2.72, p = 0.003; study 2: 2.91 vs 2.63, p < 0.001). The reported rate of a previous breast biopsy in cases was 17.8% (study 1) and 22.5% (study 2), and in controls it was 14.5% (study 1) and 15.1% (study 2). The difference in biopsy rate between cases and controls was statistically

significant in study 2 (p = 0.005), but was similar (in study 2) to the PROCAS study as a whole. In study 1 significantly fewer cases than controls reported being of "white" ethnic origin (91.3% vs 94.5%, p = 0.003), and fewer cases than controls reported having children in study 2 (85.8% vs 90.2%, p = 0.023).

In study 1, VAS results were missing for 46 cases of cancer, Quantra failed to produce results for one case and one control, Volpara failed for one case, and Densitas failed for 6 cases and 62 controls. In study 2 there were missing density results for two cases of cancer assessed by VAS, for one case and one control assessed by Quantra and for 7 cases and 34 controls assessed by Densitas.

## Study 1: screen-detected cancers

In study 1 after full adjustment, the strongest predictor of breast cancer risk was visually assessed density (Table 2, Fig. 1), with an odds ratio (OR) of 4.37 (95% CI 2.72-7.03) in the highest quintile of density compared with the lowest. When quantized in Boyd categories (Table 3), the adjusted OR of those with greater than 50% density was 6.73 (95% CI 3.64-12.45) compared to those with density 10% or lower. Volpara percent density provided the next strongest association with cancer, with an OR for the highest quintile of 2.42 (95% CI 1.56-3.78) (Table 2, Fig. 1). When guantized in Volpara Density Grades (VDG 5<sup>th</sup> edition), the OR of VDG4 was 4.39 (95% CI 2.28-8.48) compared with VDG1 (Table 3). Both visually assessed density and Volpara percent density showed a significant and clear trend with increasing density ( $\chi^2$  trend 35.6, p < 0.001 and 11.2, p < 0.001, respectively). Percent density measured by Densitas and Cumulus was also statistically significant (Table 2, Fig. 1), with ORs of 2.17 (95% CI 1.41-3.33) and 2.12 (95% CI 1.30–3.45), respectively in the highest quintile of percent density compared with the lowest, and for Quantra there was no significant association (OR = 1.02, 95% CI 0.67-1.54). The relationship with dense volume is shown in Table 2; generally associations tended to be slightly lower than those for percent density. In the subset of women with all density measures VAS was a significantly better predictor of breast cancer risk than all other methods (Table 2, Additional file 1: Table S2). The matched concordance index for VAS was 0.651 (95% CI 0.611-0.691) demonstrating better discrimination between cases and controls than all other methods (Table 4).

# Study 2: prior mammograms

In study 2 visually assessed density had the strongest association with subsequent cancer in the fully adjusted

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	Study 1					Study 2					PROCAS	
	Control sub	jects	Case subjec	ts		Control subj	ects	Case subjects			(n = 57902)	
	(n = 1098)		(n = 366)			(n = 1014)		(n = 338)				
	Number	Percent	Number	Percent	P value	Number	Percent	Number	Percent	P value	Number	Percent
Age at consent (years)												
< 50	117	10.7	40	10.9		50	4.9	17	5.0		7173	12.4
50-54	316	28.8	104	28.4		203	20.0	66	19.5		16962	29.3
55-59	157	14.3	53	14.5		179	17.7	61	18.0		11046	19.1
60-64	237	21.6	79	21.6		315	31.1	105	31.1		11223	19.4
65–69	197	17.9	99	18.0		210	20.7	70	20.7		8552	14.8
70+	74	6.7	24	6.6	1.000	57	5.6	19	5.6	1.000	2946	5.1
Median (IQR)	58	51-64	58	51-64	0.988	61	55-65	61	55-65	0.965	57	51-63
Menopausal status												
Perimenopausal	170	15.5	57	15.6		145	14.3	49	14.5		10760	18.6
Postmenopausal	747	68.0	248	67.8		769	75.8	256	75.7		37201	64.2
Premenopausal	136	12.4	46	12.6		67	6.6	22	6.5		6869	11.9
Unknown	45	4.1	15	4.1	1.000	33	3.3	11	3.3	1.000	3072	5.3
HRT use												
Never	669	63.7	250	68.3		518	51.1	177	52.4		36505	63.0
Previous	342	31.1	93	25.4		379	37.4	122	36.1		16438	28.4
Current	56	5.1	19	5.2		110	10.8	37	10.9		4421	7.6
Unknown	1	0.1	4	1.1	0.302 <sup>a</sup>	7	0.7	2	0.6	0.972	538	0.9
BMI (kg/m <sup>2</sup> )												
< 25	345	31.4	112	30.6		334	32.9	108	32.0		20774	35.8
25–29	353	32.1	122	33.3		346	34.1	118	34.9		18969	32.8
≥30	327	29.8	105	28.7		272	26.8	88	26.0		14256	24.6
Unknown	73	6.6	27	7.4	0.918	62	6.1	24	7.1	0.904	3933	6.8
Median (IQR)	27.2	24.0-31.2	27.8	24.0-30.8	0.581	26.6	23.9–30.5	26.6	23.9-30.5	0.517	26.4	23.6-30.3
Ethnic origin												
White	1038	94.5	334	91.3		924	91.1	304	6.68		52689	91.0
Other/unknown	50	4.6	32	8.7	0.003	06	8.9	34	10.1	0.511	5213	9.0
Year of mammogram												
2009	22	2.0	0	0.0							2372	4.1
2010	607	55.3	43	11.7		196	19.3	70	20.7		14761	25.5

Control subjects Care	Stu	udy 1					Study 2					PROCAS	
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	0	ontrol subjec	ts	Case subjec	ts		Control sub	jects	Case subjec	ts		(n = 57902)	
Mumber Percent Number Second	· (J)	= 1098)		(n = 366)			(n = 1014)		(n = 338)				
2011 396 36.1 130 35.5 55.4 177 52.4   2012 70 64 129 32.2 264 260 35.4 25.4   2013 3 0 32 32 32 32 56 25.4   2013 0 0 0 3 30 32 52.4 56   2013 0 0 0 10 13 10 25 66   1ntial Tyter-Cuck (troyar %) 10 0 003 263 263 27 26   1ntial Tyter-Cuck (troyar %) 218 32 219 32 219 21	Nr	umber	Percent	Number	Percent	P value	Number	Percent	Number	Percent	P value	Number	Percent
2012 70 64 129 35.2 264 256 254   2013 3 0.3 30 80 82 3 00 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90 3 90	2011 39	96	36.1	130	35.5		553	54.5	177	52.4		18350	31.7
2013 3 03 30 82 1 01 3 09   2014 0 0 33 90 3 90 2 06 00   2015 0 00 1 03 90 2 06 00   2015 0 00 1 03 20 20 00 2 06   Inital Tyrer-Cuck (10-year %) 1 2 259 205 259 205 205 2003 263	2012 70	6	6.4	129	35.2		264	26.0	86	25.4		14214	24.5
D14 0 00 33 90 01 2 06 2 06   2015 0 00 1 03 <000 <sup>1</sup> 1 20 06 00 1 03 00 2 06 01   Initi TyrerCuzick (10-year 9) 272 218 - 352 295 - 300 003 263 210 - 347 291 225 - 402 <10	2013 3		0.3	30	8.2		-	0.1	ŝ	0.9		3499	6.0
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Initial Tyrer-Cuzick (10-year %) Median (10)% 2/2 218 - 3/2 29 - 3/2 29 - 3/2 218 - 3/2 2/2 2/2 2/2 2/2 2/2 2/2 2/2 2/2 2/2	2015 0		0.0	-	0.3	< 0.001 <sup>b</sup>					0.769 <sup>c</sup>	006	1.6
Wedian (QR) 272 218 - 352 295 - 300 0003 263 210-347 291 225-402 <1   Family history 83 731 249 680 754 744 29 678   None 83 731 249 680 754 744 29 678   FBR only 147 134 51 139 107 106 47 130   SDR only 147 134 51 139 126 678 91 130   Current alcohu use 134 51 139 126 126 73 27 20 59 00   No 283 258 89 233 243 75 15 27 01 27   No 283 258 89 275 27 20 27 01   No 283 253 269 93 15 27 27 01 01	ial Tyrer-Cuzick (10-year %)												
Family history Family history 754 744 229 678   None 803 731 249 680 754 74 29 678   FDR only 104 95 42 115 107 106 47 130   FDR only 147 134 51 139 126 124 42 130   SDR only 147 134 51 139 126 124 42 130   SDR only 147 134 51 139 126 124 42 130   Current alcohol use 233 253 89 273 269 716 713 713   Vis 736 157 27 269 716 713 713   Any children 18 166 716 716 713 713 713 714   Any children 183 213 883 215 716 713 713 713<	Median (IQR) 2.7	72	2.18 – 3.52	2.95	2.29–3.90	0.003	2.63	2.10-3.47	2.91	2.25-4.02	< 0.001	2.67	2.12-3.47
None 803 731 249 680 754 744 229 678   FDR only 104 95 42 115 107 106 47 130   SDR only 147 134 51 139 166 115 124 130   SDR only 147 134 51 139 126 124 124   SDR only 147 136 24 66 0.112 27 20 59 01   Current alcohol use 233 258 89 243 273 269 373 269 375 275 01   Ves 736 155 269 716 713 713 713   Any children 1 1 1 275 15 1<1	nily history												
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FDR and SDR 44 40 24 66 0.112 27 20 59 01   Current alcohol use 233 258 89 24.3 273 26.9 93 27.5 01   No 283 258 89 73.5 776 71.6 241 71.3 27.5 01   Ves 797 72.6 269 73.5 72.6 71.6 241 71.3 01   Unknown 18 1.6 8 2.2 0.697 15 1.5 4 1.2 01   Any children 1 1 4 1.1 4 1.2 01 01   Any children 128 11.7 43 11.7 98 97 45 13.3 01 01   Ves 970 88.3 323 88.3 915 902 290 658 01 01   Ves 910 82.5 240 13	5DR only 14	17	13.4	51	13.9		126	12.4	42	12.4		7998	13.8
Current alcohol use   No 283 25.8 89 24.3 273 26.9 93 275   Ves 797 72.6 269 73.5 726 71.6 241 71.3   Ves 797 72.6 269 73.5 726 71.6 241 71.3   Unknown 18 1.6 8 2.2 0.697 15 1.5 4 1.2 0.1   Any children 1 1 43 11.7 98 97 45 133   No 128 11.7 43 11.7 983 915 902 290 858   Ves 970 883 333 88.3 1 0.1 0.1 3 0.9 01   Ves 910 883 1 0.963 1 3 0.9 01   Ves 910 82.9 28.7 0.6 28.7 0.9 0.9	<sup>-</sup> DR and SDR 44		4.0	24	9.9	0.112	27	2.7	20	5.9	0.008	1788	3.1
No 283 258 89 243 273 269 93 275   Yes 797 726 269 735 726 716 241 713   Unknown 18 1.6 8 2.2 0.697 15 4 1.2 0.5   Any children 1 1 43 11.7 43 11.7 98 97 45 133   Any children 128 11.7 43 11.7 98 97 45 133   Ves 970 883 323 883 1 0.16 3 0.9 01   Ves 970 883 323 883 1 0.1 3 0.9 01   Ves 910 883 1 0.95 1 3 0.9 01   Ves 910 821 1 0.1 1 0.1 3 0.9 01   Vior biopoy of breast 15	rrent alcohol use												
Yes797726716241713Unknown181.682.20.697151.541.20.5Any children1.20.5Any children1.20.5Any children1.20.5No128 <td>Vo 28.</td> <td>33</td> <td>25.8</td> <td>89</td> <td>24.3</td> <td></td> <td>273</td> <td>26.9</td> <td>93</td> <td>27.5</td> <td></td> <td>15815</td> <td>27.3</td>	Vo 28.	33	25.8	89	24.3		273	26.9	93	27.5		15815	27.3
Unknown 18 1.6 8 2.2 0.697 15 1.5 4 1.2 0.6   Any children 1 1 43 11.7 43 11.7 43 13.3   No 128 11.7 43 11.7 98 9.7 45 13.3   Ves 970 88.3 32.3 88.3 915 90.2 290 85.8   Unknown 1 0.963 1 0.19 3 0.9 0.1   Prior biops of breast 1 2 284 77.6 837 82.5 746   Ves 159 14.5 65 17.8 153 151 76 225	Yes 79	76	72.6	269	73.5		726	71.6	241	71.3		41130	71.0
Any children Any children 43 11.7 43 11.7 98 9.7 45 133   No 128 11.7 43 11.7 98 9.7 45 133   Yes 970 88.3 323 88.3 915 90.2 290 85.8   Unknown 1 1 0.963 1 0.1 3 0.9 0.0   Pior biopsy of breast 1 0.963 1 0.1 3 0.9 0.0   No 910 82.9 284 77.6 83.7 82.5 746   Yes 159 14.5 65 17.8 153 15.1 76 225	Jnknown 18	~	1.6	œ	2.2	0.697	15	1.5	4	1.2	0.908	957	1.7
No 128 11.7 43 11.7 98 9.7 45 133   Yes 970 88.3 323 88.3 915 90.2 290 858   Unknown 1 2 1 0.963 1 0.1 3 0.9 0.0   Prior biops of breast 1 0.963 1 0.1 3 0.9 0.0   No 910 82.9 284 776 837 82.5 252 746   Ves 159 14.5 65 178 153 15.1 76 225	y children												
Yes 970 88.3 233 88.3 915 90.2 290 85.8   Unknown Unknown 0.963 1 0.1 3 0.9 0.0   Prior biopsy of breast   0.963 1 0.1 3 0.9 0.1   No 910 82.9 284 77.6 837 82.5 74.6   Yes 159 14.5 65 17.8 15.1 76 22.5	No 12,	8	11.7	43	11.7		98	9.7	45	13.3		7384	12.7
Unknown 0.963 1 0.1 3 0.9 0.0   Prior biopsy of breast No 910 82.9 284 77.6 83.7 82.5 74.6   Ves 159 14.5 65 17.8 15.1 76 22.5	Yes 97,	0,	88.3	323	88.3		915	90.2	290	85.8		50411	87.1
Prior biopsy of breast No 910 82.9 284 77.6 837 82.5 252 74.6   Ves 159 14.5 65 17.8 15.1 76 22.5	Jnknown					0.963		0.1	С	0.9	0.023 <sup>d</sup>	107	0.2
No 910 82.9 284 77.6 837 82.5 252 746   Yes 159 14.5 65 17.8 153 15.1 76 22.5	or biopsy of breast												
Yes 159 14.5 65 17.8 153 15.1 76 22.5	Vo 91,	0	82.9	284	77.6		837	82.5	252	74.6		47359	81.8
	Yes 15	6	14.5	65	17.8		153	15.1	76	22.5		8911	15.4
Unknown 29 2.6 17 4.6 0.041 24 2.4 10 3.0 0.	Jnknown 29	~	2.6	17	4.6	0.041	24	2.4	10	3.0	0.005	1632	2.8

<sup>\*</sup>In study 1 cases are women with breast cancer detected at first screen on entry to the PROCAS study and in study 2 cases are women veubsequent screen or between screening rounds *IOR* interquartile range, *HRT* hormone replacement therapy, *BMI* body mass index, *FDR* first-degree relative, *SDR* second-degree relative <sup>a</sup>Excludes unknown <sup>b</sup>2013-2015 combined <sup>c</sup>2012-2014 combined <sup>d</sup>Unknown combined with no children

Table 2 Ris	sk of developing	breast cancer by	density measures	s (highest versus the	lowest quintile <sup>a</sup> (referent))
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					Subset	with data for all m	ethods <sup>b</sup>	
	Univaria	ate	Adjuste	dc	Univaria	ate	Adjuste	d <sup>c</sup>
	OR	95% CI	OR	95% CI	OR	95% CI	OR	95% CI
Study 1								
	1.00	(referent)	1.00	(referent)	1.00	(referent)	1.00	(referent)
VAS (%)	4.45	(2.77–7.15)	4.37	(2.72–7.03)	5.61	(3.29–9.56)	5.44	(3.18–9.29)
Volpara gland volume (cm <sup>3</sup> )	2.13	(1.40–3.24)	2.09	(1.37–3.18)	2.00	(1.24–3.21)	1.97	(1.22–3.18)
Volpara breast density (%)	2.44	(1.57–3.80)	2.42	(1.56–3.78)	2.38	(1.45–3.91)	2.41	(1.46–3.97)
Cumulus dense area (cm <sup>2</sup> )	2.11	(1.32–3.38)	2.15	(1.34–3.45)	2.08	(1.28–3.27)	2.12	(1.30–3.45)
Cumulus breast density (%)	2.09	(1.29–3.40)	2.12	(1.30–3.45)	2.20	(1.32–3.64)	2.23	(1.34–3.71)
Quantra gland volume (cm <sup>3</sup> )	0.86	(0.58–1.30)	0.83	(0.55–1.25)	0.72	(0.45–1.15)	0.71	(0.44–1.13)
Quantra breast density (%)	1.05	(0.70–1.59)	1.02	(0.67–1.54)	1.11	(0.70–1.77)	1.08	(0.68–1.72)
Densitas dense area (cm <sup>2</sup> )	1.44	(0.96–2.16)	1.41	(0.93–2.12)	1.62	(1.01–2.59)	1.61	(1.00–2.58)
Densitas breast density (%)	2.30	(1.50–3.52)	2.17	(1.41–3.33)	2.19	(1.35–3.56)	2.10	(1.29–3.41)
Study 2								
	1.00	(referent)	1.00	(referent)	1.00	(referent)	1.00	(referent)
VAS (%)	4.54	(2.86–7.22)	4.48	(2.79–7.18)	4.41	(2.76–7.06)	4.36	(2.70–7.04)
Volpara gland volume (cm <sup>3</sup> )	2.72	(1.79–4.14)	2.66	(1.74–4.08)	2.71	(1.77–4.14)	2.65	(1.72–4.09)
Volpara breast density (%)	2.78	(1.74–4.44)	2.87	(1.77–4.64)	2.61	(1.62–4.19)	2.71	(1.67–4.39)
Quantra gland volume (cm <sup>3</sup> )	1.36	(0.90–2.06)	1.28	(0.84–1.95)	1.32	(0.87–2.01)	1.24	(0.80–1.90)
Quantra breast density (%)	1.32	(0.86–2.03)	1.32	(0.85–2.05)	1.30	(0.84–2.00)	1.32	(0.85–2.05)
Densitas dense area (cm <sup>2</sup> )	2.34	(1.56–3.52)	2.23	(1.48–3.38)	2.29	(1.50–3.50)	2.14	(1.40–3.29)
Densitas breast density (%)	2.45	(1.57–3.82)	2.34	(1.50–3.68)	2.44	(1.56–3.80)	2.34	(1.49–3.66)

OR odds ratio, CI confidence interval, VAS visual analogue scale, results in italics indicate statistically significant results (p<0.05)

<sup>a</sup>Quintiles based on distribution amongst controls

<sup>b</sup>Study 1: 239 cases with 3 controls, 62 with 2 controls and 2 with 1 control; study 2: 296 cases with 3 controls, 31 with 2 controls and 2 with 1 control <sup>c</sup>Adjusted for Tyrer-Cuzick score; study 2 also adjusted for parity

Figures in italics denote statistically significant results p < 0.01

models, with an OR of 4.48 (95% CI 2.79-7.18) in the highest quintile of density compared with the lowest (Table 2). When quantized in Boyd categories (Table 3), the OR of those with density > 50% was 5.45 (95% CI 3.00–9.89) compared to those with density  $\leq 10\%$ . Volpara percent density had the next strongest association with cancer, with an OR for the highest quintile of 2.87 (95% CI 1.77-4.64) (Table 2, Fig. 2). When quantized in Volpara Density Grades (5<sup>th</sup> Edition), the OR of VDG4 was 3.00 (95% CI 1.54-5.86) compared with VDG1 (Table 3). Both visually assessed density and Volpara percent density showed a dose response relationship with increasing density ( $\chi^2$  trend 42.7, p < 0.001 and 13.8, p < 0.001, respectively). For Densitas and Quantra, those with percentage density in the highest quintile had ORs of 2.34 (95% CI 1.50-3.68) and 1.32 (95% CI 0.85-2.05), respectively (Table 2, Fig. 2). VAS predicted breast cancer risk significantly better than all other density methods in the subset of women who had density measured by all four methods (Table 2, Additional file 1: Table S2). The matched concordance index for VAS was

0.647 (95% CI 0.607–0.688) demonstrating better discrimination between cases and controls than all other methods (Table 4).

## Discussion

Visual assessment of breast density recorded on a VAS was the strongest predictor of breast cancer risk, both in the contralateral breast of women with screen-detected cancers and in the average of bilateral mammographic views prior to the detection of cancer. It is unlikely that the presence of cancer influenced visual assessment in study 1, since a blinded re-read of images from the contralateral breast by four readers showed no evidence of bias [23] and the ORs were similar to those in study 2. There is strong association between the VAS and breast cancer despite known inter-observer variability [32]; since the average VAS score of two readers was used it is likely that cases falling into the top and bottom quintiles of density do so unequivocally.

Volpara and Densitas percent density had the next strongest associations with cancer in both studies, with



categorisation into VDG having the largest odds ratio. Volpara, Quantra and Cumulus did not have as strong an association with breast cancer in study 1 as previously reported [13]. This may be due to differences in the approach used; Eng et al. analysed 414 cases from one hospital and 685 unmatched controls from a screening service based in London, adjusting for age, BMI and reproductive variables in the analysis, whilst we analysed 366 cases using Volpara and Quantra, and 311 using Cumulus, with 3 well-matched controls per case all recruited from the same screening programme. There were also a number of differences between the study populations, with our study population tending to be younger, with more women of white ethnicity and with higher BMI and being less likely to be postmenopausal and to have had children. Density distributions also differed across the two studies, with the current study having lower median (IQR) percent density assessed by Volpara (4.9, 3.5-7.4) and Quantra (11, 8-14), but higher percent density for Cumulus (20.3, 11.6-30.3) [13]. Our version of Volpara was later (1.5.0 vs 1.0) and we applied a Volpara macro for outlier rejection; our version of Quantra was also more recent (2.0 vs 1.3). For Cumulus, the difference might be due to reader experience.

Study 2 examined the relationship between mammographic density in mammograms prior to the detection of cancer, and in matched controls that subsequently remained cancer free. This enables us to evaluate which mammographic density methods are most appropriate for stratifying women attending breast screening. Whilst visual assessment was most strongly associated with cancer, it is unlikely to be used widely for population-based stratified screening; we conclude that Volpara or Densitas percentage density provide a pragmatic solution. However, we hypothesise that methods that measure purely the quantity or relative proportion of dense tissue do not fully capture the mammographic risk in the same way as visual assessment by experts, who can see not only the quantity of dense tissue but the location and pattern. The addition of algorithms that automatically quantify mammographic pattern to automated density software could potentially provide a solution that more closely reproduces visual assessment. Recent research in

	5	2 m) cure		2												
									Subset	with data for	all metho	ds <sup>b</sup>				
	Contro	slo	Cases		Univariá	ate	Adjuste	,d <sup>a</sup>	Control	S	Cases		Univaria	ite	Adjuste	da
	2	Percent	Z	Percent	Ю	95% CI	OR	95% CI	Z	Percent	Z	Percent	OR	95% CI	OR	95% CI
Study 1																
VAS (mean) - Boyc	ł categorie	s (combining	> 50-75%	and > 75%)												
≥ 0-10%	208	(18.9)	29	(9.1)	1.00	(referent)	1.00	(referent)	176	(21.1)	25	(8.3)	1.00	(referent)	1.00	(referent)
> 10-25%	436	(39.7)	115	(35.9)	2.17	(1.37–3.42)	2.16	(1.37–3.40)	331	(39.7)	111	(36.9)	2.67	(1.63–4.38)	2.65	(1.62–4.34)
> 25-50%	379	(34.5)	133	(41.6)	3.25	(2.05–5.18)	3.20	(2.01–5.09)	277	(33.3)	125	(41.5)	3.95	(2.38–6.54)	3.84	(2.31–6.36)
> 50%	75	(6.8)	43	(13.4)	6.92	(3.74–12.79)	6.73	(3.64–12.45)	49	(5.9)	40	(13.3)	8.30	(4.28–16.07)	8.02	(4.13–15.56)
Volpara Density Gı	ades 4th E	dition														
1 (<4.5)	496	(45.2)	124	(34.0)	1.00	(referent)	1.00	(referent)	400	(48.0)	108	(35.9)	1.00	(referent)	1.00	(referent)
2 (4.5-<7.5)	339	(30.9)	131	(35.9)	1.78	(1.31–2.41)	1.74	(1.29–2.37)	258	(31.0)	113	(37.5)	1.90	(1.35–2.66)	1.88	(1.34–2.63)
3 (7.5-<15.5)	229	(20.9)	85	(23.3)	1.85	(1.29–2.64)	1.86	(1.30–2.68)	153	(18.4)	64	(21.3)	1.92	(1.28–2.90)	1.94	(1.29–2.94)
4 (15.5+)	34	(3.1)	25	(6.8)	4.18	(2.24–7.77)	4.07	(2.18–7.60)	22	(2.6)	16	(5.3)	3.73	(1.80–7.75)	3.58	(1.72–7.44)
Volpara Density Gr	ades 5th E	dition														
1 (< 3.5)	274	(25.0)	62	(17.0)	1.00	(referent)	1.00	(referent)	216	(25.9)	50	(16.6)	1.00	(referent)	1.00	(referent)
2 (3.5-<7.5)	561	(51.1)	193	(52.9)	1.68	(1.20–2.34)	1.68	(1.20–2.35)	442	(53.1)	171	(56.8)	1.88	(1.29–2.73)	1.89	(1.30–2.75)
3 (7.5-<15.5)	229	(20.9)	85	(23.3)	2.00	(1.32–3.02)	2.04	(1.34–3.09)	153	(18.4)	64	(21.3)	2.22	(1.38–3.57)	2.27	(1.41–3.66)
4 (15.5+)	34	(3.1)	25	(6.8)	4.44	(2.31–8.54)	4.39	(2.28–8.48)	22	(2.6)	16	(5.3)	4.26	(1.98–9.17)	4.14	(1.92–8.95)
Quantized density (B	IRADS)															
1	54	(4.9)	27	(7.4)	1.00	(referent)	1.00	(referent)	47	(2.6)	21	(7.0)	1.00	(referent)	1.00	(referent)
2	682	(62.2)	220	(60.3)	0.63	(0.39–1.02)	0.61	(0.38–0.99)	530	(63.6)	193	(64.1)	0.85	(0.50-1.44)	0.80	(0.47-1.37)
m	317	(28.9)	102	(27.9)	0.63	(0.37-1.05)	0.58	(0.34–0.99)	226	(27.1)	75	(24.9)	0.77	(0.43-1.38)	0.71	(0.39–1.28)
4	44	(4.0)	16	(4.4)	0.71	(0.33-1.52)	0.62	(0.29–1.33)	30	(3.6)	12	(4.0)	0.92	(0.39–2.17)	0.79	(0.33–1.88)
Densitas: BIRADS 4	_															
0-24	387	(37.4)	105	(35.2)	1.00	(referent)	1.00	(referent)	326	(39.1)	94	(31.2)	1.00	(referent)	1.00	(referent)
25–50	599	(57.8)	219	(60.8)	1.53	(1.15–2.04)	1.48	(1.11-1.98)	470	(56.4)	186	(61.8)	1.53	(1.12–2.08)	1.48	(1.09–2.03)
51+	50	(4.8)	36	(10.0)	3.46	(2.04–5.87)	3.21	(1.88–5.47)	35	(4.2)	21	(0.7)	2.47	(1.34–4.55)	2.33	(1.26–4.30)
Densitas: BIRADS 5	10															
∢	194	(18.7)	46	(12.8)	1.00	(referent)	1.00	(referent)	171	(20.5)	37	(12.3)	1.00	(referent)	1.00	(referent)
Β	552	(53.3)	191	(53.1)	1.58	(1.08–2.30)	1.50	(1.03–2.19)	446	(53.5)	174	(57.8)	1.97	(1.29–2.99)	1.87	(1.23–2.85)
U	278	(26.8)	117	(32.5)	2.01	(1.33–3.04)	1.92	(1.27–2.91)	206	(24.7)	85	(28.2)	2.12	(1.34–3.37)	2.03	(1.28–3.23)
D	12	(1.2)	9	(1.7)	2.79	(0.94–8.24)	2.50	(0.84–7.48)	00	(1.0)	5	(1.7)	3.71	(1.09–12.66)	3.30	(0.95–11.42)

Table 3 Density measures by categorical scales

									Subset	with data for	all metho	dsb				
	Contro	ls	Cases		Univari	ate	Adjuste	e pe	Contro	ls	Cases		Univarià	ate	Adjuste	ed <sup>a</sup>
	z	Percent	2	Percent	SOR	95% CI	OR	95% CI	Z	Percent	z	Percent	OR	95% CI	OR	95% CI
Study 2																
VAS (mean) - Boyc	l categorie	s (combining	>50-75%	i and >75%)												
≥ 0-10%	193	(19.0)	34	(10.1)	1.00	(referent)	1.00	(referent)	180	(18.7)	32	(6.7)	1.00	(referent)	1.00	(referent)
> 10-25%	401	(39.5)	103	(30.5)	1.61	(1.05–2.49)	1.55	(1.00–2.40)	383	(39.8)	100	(30.4)	1.61	(1.03–2.50)	1.55	(0.99–2.42)
> 25-50%	356	(35.1)	154	(45.6)	2.92	(1.89–4.52)	2.87	(1.85–4.46)	329	(34.2)	152	(46.2)	3.02	(1.94–4.71)	2.97	(1.90–4.67)
> 50%	64	(6.3)	45	(13.3)	5.52	(3.08–9.91)	5.45	(3.00–9.89)	60	(6.2)	45	(13.7)	5.65	(3.12–10.23)	5.63	(3.07–10.3)
Volpara Density Gr	ades 4th E	dition														
1 (< 4.5)	474	(46.7)	126	(37.3)	1.00	(referent)	1.00	(referent)	447	(46.5)	123	(37.4)	1.00	(referent)	1.00	(referent)
2 (4.5- < 7.5)	336	(33.1)	120	(35.5)	1.45	(1.07–1.96)	1.44	(1.06–1.95)	315	(32.7)	118	(35.9)	1.44	(1.06–1.95)	1.43	(1.05–1.95)
3 (7.5-<15.5)	167	(16.5)	73	(21.6)	1.96	(1.34–2.87)	2.08	(1.40–3.08)	156	(16.2)	69	(21.0)	1.81	(1.23–2.67)	1.92	(1.29–2.86)
4 (15.5+)	37	(3.6)	19	(5.6)	2.47	(1.30–4.70)	2.61	(1.34 - 5.10)	34	(3.5)	19	(5.8)	2.56	(1.33–4.91)	2.73	(1.38–5.38)
Volpara Density Gr	ades 5th E	dition														
1 (<3.5)	205	(20.2)	47	(13.9)	1.00	(referent)	1.00	(referent)	195	(20.3)	47	(14.3)	1.00	(referent)	1.00	(referent)
2 (3.5-<7.5)	570	(56.2)	189	(55.9)	1.55	(1.07–2.24)	1.53	(1.06–2.23)	533	(55.4)	184	(55.9)	1.51	(1.04–2.19)	1.51	(1.04–2.21)
3 (7.5-<15.5)	193	(19.0)	78	(23.1)	2.09	(1.33–3.28)	2.16	(1.36–3.43)	181	(18.8)	74	(22.5)	1.92	(1.22–3.04)	1.99	(1.25–3.17)
4 (15.5+)	46	(4.5)	24	(7.1)	2.92	(1.53–5.58)	3.00	(1.54–5.86)	43	(4.5)	24	(7.3)	2.82	(1.47–5.40)	2.91	(1.48–5.72)
Quantized density	(BIRADS)															
-	32	(3.2)	12	(3.6)	1.00	(referent)	1.00	(referent)	32	(3.3)	12	(3.6)	1.00	(referent)	1.00	(referent)
2	654	(64.5)	197	(58.5)	0.81	(0.39–1.70)	0.87	(0.42-1.82)	616	(64.0)	192	(58.4)	0.82	(0.39–1.71)	0.88	(0.42–1.83)
3	287	(28.3)	108	(32.0)	1.05	(0.49–2.27)	1.12	(0.52-2.41)	267	(27.8)	105	(31.9)	1.04	(0.48–2.23)	1.11	(0.52–2.39)
4	40	(3.9)	20	(5.9)	1.51	(0.59–3.88)	1.54	(0.59-4.01)	37	(3.8)	20	(6.1)	1.50	(0.58-3.87)	1.54	(0.59-4.02)
Densitas: BIRADS 4	_*															
0-24	354	(36.1)	95	(28.7)	1.00	(referent)	1.00	(referent)	345	(35.9)	95	(28.9)	1.00	(referent)	1.00	(referent)
25-50	586	(59.8)	208	(62.8)	1.37	(1.02–1.83)	1.34	(1.00–1.80)	568	(29.0)	207	(62.9)	1.37	(1.02–1.83)	1.34	(1.00–1.80)
51+	40	(4.1)	28	(8.5)	2.94	(1.65–5.24)	2.83	(1.55–5.16)	39	(4.1)	27	(8.2)	2.86	(1.60–5.11)	2.76	(1.51–5.05)
Densitas: BIRADS 5	1.5															
A	96	(8.8)	25	(2.6)	1.00	(referent)	1.00	(referent)	94	(8.6)	25	(7.6)	1.00	(referent)	1.00	(referent)
В	574	(58.6)	185	(57.9)	1.25	(0.77-2.01)	1.20	(0.74-1.94)	558	(58.0)	185	(56.2)	1.25	(0.77-2.01)	1.20	(0.74–1.94)
U	294	(30.0)	110	(33.2)	1.47	(0.88-2.45)	1.46	(0.87–2.45)	284	(29.5)	108	(32.8)	1.46	(0.88–2.44)	1.45	(0.87-2.44)
D	16	(1.6)	11	(3.3)	2.90	(1.14–7.42)	2.59	(0.99–6.78)	16	(1.7)	11	(3.3)	2.90	(1.13–7.41)	2.58	(0.99–6.76)
N number, OR odds	ratio, C/ c	onfidence in	terval, VA	S visual anald	ogue scale	, BIRADS Breast I	Imaging F	Reporting and Di	ata Systen	n, results in i	talics indi	cate statistic	ally signifi	icant results (p<(	0.05)	

<sup>a</sup>Adjusted for Tyrer-Cuzick score; study 2 also adjusted for parity <sup>b</sup>Study 1: 239 cases with 3 controls, 62 with 2 controls and 2 with 1 control; study 2: 296 cases with 3 controls, 31 with 2 controls and 2 with 1 control

## Table 4 Matched concordance index (mC)

	Study	1		Study	2	
	mC <sup>a</sup>	95% C	l	тС <sup>ь</sup>	95% C	1
VAS (%)	0.651	0.611	0.691	0.647	0.607	0.688
Volpara breast density (%)	0.571	0.528	0.618	0.575	0.534	0.615
Volpara gland volume (cm <sup>3</sup> )	0.553	0.513	0.591	0.586	0.546	0.627
Quantra breast density (%)	0.510	0.469	0.552	0.543	0.504	0.584
Quantra gland volume (cm <sup>3</sup> )	0.487	0.447	0.528	0.531	0.490	0.574
Densitas breast density (%)	0.571	0.526	0.612	0.587	0.548	0.628
Densitas dense area (cm <sup>2</sup> )	0.535	0.496	0.574	0.577	0.537	0.616
Cumulus breast density (%)	0.582	0.541	0.623	-	-	-
Cumulus dense area (cm <sup>2</sup> )	0.558	0.516	0.599	-	-	-

CI confidence interval, VAS visual analogue scale

<sup>a</sup>Study 1: 239 cases with 3 controls, 62 with 2 controls, 2 with 1 control <sup>b</sup>Study 2: 296 with 3 controls, 31 with 2 controls, 2 with 1 control

this area has proved promising [33–36], although there is as yet no consensus as to the best method of encapsulating texture information within risk assessment.

## Strengths and limitations

The strengths of this study include the ability to assess the relationship between several measures of mammographic density and risk of breast cancer. As well as examining the association between mammographic density and breast cancer risk, we were able to establish the temporal relationship in study 2. We also gathered detailed information in relation to a number of covariates (demographic, hormonal, reproductive, lifestyle and family history) via a self-reported questionnaire at entry to PROCAS [20]. Uptake to PROCAS was relatively low (38%), which may have biased the population to those with higher or lower risk, for example, the proportion of women in the PROCAS study who were overweight or obese was significantly lower than in the general population of Greater Manchester [37]. In addition, in study 1, due to the nature of the study design, whereby controls had to have had a subsequent cancer-free mammogram after entry to the PROCAS study, the year of mammogram in controls tended to be earlier than in cases, this may have had an impact on density measures due to changes in mammography technology, and the use of different mammographic machines over time.

# Conclusions

Visual assessment of density, recorded on a VAS and averaged between two independent readers, is a strong predictor of breast cancer risk both in mammograms taken before the detection of cancer and in images of the opposite breast at the time of detection. Percentage density measured by Volpara and Densitas also showed a strong association with breast cancer risk amongst the automated measures evaluated, providing practical automated methods for risk stratification in personalized screening programmes.



# **Additional file**

Additional file 1: Table S1. Risk of developing breast cancer using continuous measures of different density methods (OR per SD). Table S2. *P* values based on likelihood ratio comparing different models for density methods using the subset of those with data on all methods. (DOCX 26 kb)

#### Abbreviations

BIRADS: Breast Imaging Reporting And Data System; BMI: Body mass index; CI: Confidence interval; FFDM: Full- field digital mammography; HRT: Hormone replacement therapy; IQR: Interquartile range; ICC: Intraclass correlation coefficient; MLO: Mediolateral oblique; OR: Odds ratio; PROCAS: Predicting Risk of Cancer At Screening Study; VAS: Visual analogue scale; VDG: Volpara Density Grade

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#### Availability of data and materials

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

## Authors' contributions

SMA conceived and designed the study, analysed and interpreted the data and drafted the manuscript. EFH performed the statistical analysis, interpreted the data and drafted the manuscript. JCS carried out the cumulus density assessment after training from JW. ARB participated in the statistical analysis. PS made substantial contributions to acquisition of questionnaire data. MW, UB, SG, YL, AJ, SB and NB made substantial contributions to acquisition of density data, each reading more than 4000 mammograms. VR was responsible for mammography reporting. RW, DGRE, AH and JC conceived and designed the study, analysed and interpreted the data, and helped to draft the manuscript. All authors were involved in critically revising the paper for intellectual content, and all read and approved the final manuscript.

#### Ethics approval and consent to participate

Ethics approval for the study was through the North Manchester Research Ethics Committee (09/H1008/81). Informed consent was obtained from all participants on entry to the PROCAS study.

#### Consent for publication

Not applicable.

#### **Competing interests**

Software licences for Volpara, Quantra and Densitas were provided free of charge under a research agreement by Volpara Health Technologies (Wellington, New Zealand), Hologic Inc (Marlborough, MA, USA) and Densitas Inc (Halifax, NS, Canada) respectively. The authors declare that they have no competing interests.

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