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Genetic predisposition to ductal carcinoma *in situ* of the breast

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Abstract

Background: Ductal carcinoma *in situ* (DCIS) is a non-invasive form of breast cancer. It is often associated with invasive ductal carcinoma (IDC), and is considered to be a non-obligate precursor of IDC. It is not clear to what extent these two forms of cancer share low-risk susceptibility loci, or whether there are differences in the strength of association for shared loci.

Methods: To identify genetic polymorphisms that predispose to DCIS, we pooled data from 38 studies comprising 5,067 cases of DCIS, 24,584 cases of IDC and 37,467 controls, all genotyped using the iCOGS chip.

Results: Most (67 %) of the 76 known breast cancer predisposition loci showed an association with DCIS in the same direction as previously reported for invasive breast cancer. Case-only analysis showed no evidence for differences between associations for IDC and DCIS after considering multiple testing.

Analysis by estrogen receptor (ER) status confirmed that loci associated with ER positive IDC were also associated with ER positive DCIS. Analysis of DCIS by grade suggested that two independent SNPs at 11q13.3 near *CCND1* (Continued on next page)

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were specific to low/intermediate grade DCIS (rs75915166, rs554219). These associations with grade remained after adjusting for ER status and were also found in IDC.

We found no novel DCIS-specific loci at a genome wide significance level of $P < 5.0 \times 10^{-8}$.

Conclusion: In conclusion, this study provides the strongest evidence to date of a shared genetic susceptibility for IDC and DCIS. Studies with larger numbers of DCIS are needed to determine if IDC or DCIS specific loci exist.

Keywords: Ductal carcinoma in situ, Association study, Genetic predisposition, Common variants

Background

Ductal carcinoma in situ (DCIS) is a non-obligate precursor of invasive breast cancer including invasive ductal/no special type carcinomas (IDC). Since the introduction of screening mammography there has been a 7-fold increase in reported DCIS incidence in the USA, primarily in postmenopausal women [1], with about 20 % of screen-detected tumors being DCIS [2]. Approximately 45–78 % of all invasive breast cancers are associated with DCIS [3, 4]. It is hypothesized in the majority of these cases that the invasive component has arisen from the DCIS as they generally share the same somatic genetic changes. The proportion of IDC associated with DCIS varies depending on subtype, with luminal and human epidermal growth factor receptor 2 (HER2)-positive IDC having more frequent DCIS (53 % and 63 %, respectively) than invasive basal breast cancers (33 %) [5].

As most DCIS is treated surgically, the natural progression of untreated DCIS is not known. However, in one small study of patients with predominantly lowgrade DCIS misdiagnosed as benign breast disease and who received no surgical intervention, 6 out of 13 patients developed ipsilateral invasive carcinoma with mean time to the development of invasive carcinoma being 9.0 years [6]. In two specific DCIS trials in which DCIS was treated with breast-conserving surgery alone with no radiotherapy, long-term follow up shows that up to 30 % of women develop a recurrence (half of which will be DCIS and half invasive cancer) by 10 years [7].

Methods for accurately predicting the behavior of DCIS are poor [8]. Although grade has not been shown to be a good predictor of recurrence many clinicians use this classification to determine the use of radiotherapy following breast-conserving surgery. There is a strong correlation between the grade of the *in situ* and co-existing invasive components in IDC, suggesting that DCIS does not progress from low through to high grade before becoming invasive [9, 10].

Most non-genetic risk factors for breast cancer have similar associations with DCIS and IDC, supporting the notion that DCIS is a precursor of invasive cancer [11, 12]. There is also evidence from epidemiological studies that there is an inherited predisposition to DCIS. Women with DCIS have been shown to be 2.4 times (95 % CI 0.8, 7.2) more likely to have an affected mother and sister with breast cancer than controls [13]. Furthermore, there is evidence from a study of almost 40,000 women that the familial relative risk of DCIS is greater than that of invasive breast cancer. For women aged 30–49 years with a family history of breast cancer the odds ratio (OR) for developing DCIS was calculated as 2.4 (95 % CI 1.1, 4.9) compared to 1.7 (95 % CI 0.9, 3.4) for invasive cancer. For women aged 50 years and above the risks were slightly reduced, but still higher for DCIS (OR = 2.2, 95 % CI 1.0, 4.2) than invasive disease (OR = 1.5, 95 % CI 1.0, 2.2) [14]. However, this was not confirmed in the Million Women Study, in which the association with family history was similar for DCIS and IDC [12].

A small part of this inherited predisposition is explained by *BRCA1/2* mutations, as mutations in these genes are found in a similar proportion of DCIS and invasive breast cancer cases [15]. For low-risk common breast cancer predisposition alleles most of the initial breast cancer association studies have not been powered to identify associations with DCIS, so it is not clear whether all the low-risk susceptibility loci that have been identified are associated with DCIS and what the strength of any associations are.

It is now evident that some low-risk susceptibility loci are associated with different pathological subtypes of breast cancer and support the hypothesis that breast tumor subtypes arise through distinct molecular pathways [16-18]. In order to identify further low-risk susceptibility loci, it will be necessary to look at specific morphological subtypes including DCIS and the cytonuclear grade and estrogen receptor (ER) status of the disease. In this study we analyzed 3,078 cases of pure DCIS collected through the ICICLE study (a study to Investigate the genetics of In situ Carcinoma of the ductaL subtype) and performed a meta-analysis with 2,352 in situ cases collected through the Breast Cancer Association Consortium (BCAC). Our aims were to assess whether any of the known low-risk breast susceptibility alleles have different associations for DCIS and IDC, and to identify if there are any DCIS-specific low-risk alleles.

Methods

Ethics statement

All studies were performed with ethical committee approval (listed in acknowledgements) and subjects participated in the studies after providing informed consent.

Study populations

Cases came from ICICLE (MREC 08/H0502/4), a UK study of DCIS, and from 37 studies forming part of the BCAC included in the Collaborative Oncological Gene-Environment Study (COGS) [19] (Additional file 1). The ICICLE study recruited patients from participating centers throughout the UK with the aim of identifying predisposition genes for DCIS. Patients aged 60 years or less at the time of diagnosis, with a current or past history of DCIS (without invasive disease of any histological subtype) were eligible. A total of 3,078 subjects were recruited following identification from local pathology reports in 97 UK hospitals. All cases were genotyped with the iCOGS chip and compared to 5,000 UK controls selected from four UK studies (BBCS 1,231 controls, SBCS 704 controls, UKBGS 370 controls, SEARCH 2,695 controls) participating in BCAC (Additional file 2) and already typed on the iCOGS chip. Controls were randomly selected prior to analysis, and were excluded from casecontrol comparisons with BCAC cases from the originating study. After excluding individuals based on genotyping quality (see subsection "Genotyping and analysis") and non-European ancestry, data for the ICICLE study available for analysis included 2,715 subjects with DCIS (cases) and 4,813 controls.

Women with all types of breast cancer were recruited into the BCAC studies. Pathological information in BCAC was collected in the individual studies but was also combined and checked through standardized data control in a central database. A total of 2,352 subjects with DCIS were identified in the central BCAC pathology database (see Additional file 3 for number of cases by study). Controls came from the 37 BCAC studies (37,654 in total).

Genotyping and analysis

After DNA extraction from peripheral blood, ICICLE samples were genotyped on the iCOGS custom Illumina iSelect array (Illumina, San Diego, CA), which contains 211,155 single nucleotide polymorphisms (SNPs), at King's College London. The remaining cases and controls were genotyped as part of the COGS project described in detail elsewhere [19]. The ICICLE cases were analyzed using the same quality control (QC) criteria as the COGS project. Briefly, genotypes were called using Illumina's proprietary GenCall algorithm and 10,000 SNPs were manually inspected to verify the algorithm calls. Individuals were excluded if genotypically non-European or not female, or had an overall call rate <95 %. SNPs were excluded with a Gen-Train score <0.4, call rate <95 % (call rate <99 % if minor allele frequency (MAF) was <0.1) and Hardy Weinberg equilibrium (HWE) value of $P < 10^{-7}$ or evidence of poor clustering on inspection of cluster plots. All SNPs with MAF <0.01 were excluded. A cryptic relatedness analysis of the whole dataset was performed using 46,789 uncorrelated SNPs and led to the exclusion of 28 cases and 18 controls due to relatedness between the ICICLE and BCAC samples (PIHAT >0.1875).

For ICICLE cases and controls, principal component analysis (PCA) was carried out on a subset of 46,789 uncorrelated SNPs and individuals or groups distinct from the main cluster (327 cases and 164 controls) were excluded using the first five principal components (PCs) (Additional file 4). Following removal of outliers, the PCA was repeated and the first five PCs were included as covariates in the analysis.

The adequacy of the case–control matching was evaluated using quantile-quantile plots of test statistics and the inflation factor (λ) calculated using 37,289 uncorrelated SNPs that were not selected by BCAC and were not within one of the four common fine-mapping regions, to minimize selection for SNPs associated with breast cancer (Additional file 5). As the majority of the SNPs on the iCOGS array are associated with breast, ovarian or prostate cancer, the SNPs selected for this analysis were taken from the set of prostate cancer SNPs, with the assumption that these SNPs were more likely to be representative of common SNPs in terms of population structure in our study.

For each SNP, we estimated a per-allele OR and reported corresponding 95 % CI using logistic regression analysis, including the five PCs as covariates, using PLINK v1.07 (http://pngu.mgh.harvard.edu/~purcell/plink/).

Genotyping and analysis of BCAC studies have been described in detail elsewhere [19]. In brief, data were analyzed using the Genotype Library and Utilities (GLU) package to estimate per-allele ORs for each SNP using unconditional logistic regression. All analyses were performed in subjects of European ancestry (determined by PC analyses) and adjusted for study and seven principal components.

Case-control ORs for DCIS cases vs controls from BCAC and ICICLE were combined using inverse varianceweighted fixed-effects meta-analysis, as implemented in METAL [20]. Case-only analyses were also carried out to compare genotype frequencies for (1) ER-positive (ER+) vs ER-negative (ER-) DCIS, (2) high grade DCIS vs low and intermediate grade DCIS, and (3) DCIS vs IDC (see Additional file 3 for number of cases by study), (4) DCIS diagnosis in patients <50 years of age vs DCIS diagnosis in patients \geq 50 years, and were used as a test for heterogeneity of ORs by tumor subtype/age (see Additional file 6 for number of cases by group). Only studies with data on both subtypes contributed to case-only analysis comparing these subtypes. Similar case-only analyses were performed for the IDC cases in these studies to assess whether any heterogeneity evident in DCIS also occurred in IDC.

Novel SNPs showing the strongest evidence of association with DCIS ($P < 6 \times 10^{-6}$) in the meta-analysis (after excluding previously reported loci) were genotyped in a phase II analysis at LGC Genomics (LGC, Teddington, UK). The phase II samples consisted of 653 DCIS cases from the ICICLE and Breakthrough Generation Studies and 1,882 controls from the ICICLE study not previously genotyped on the iCOGS chip. All individuals included in the analysis were of European ancestry (self-reported).

For the known breast cancer predisposition loci P <0.00066 was considered statistically significant (with Bonferroni correction for multiple testing on 76 known loci). All of the known breast cancer susceptibility loci were included in the iCOGS chip with the exception of rs2284378 (20q11), which was identified as an ER–breast cancer predisposition SNP after the iCOGS chip was developed [21].

Assessment of grade and ER status

For the ICICLE study, information on cytonuclear grade of DCIS was available for 2,578 cases, mostly from the local histopathology reports. In 200 cases where the grade data were missing from the report but the tumor block was available, an H&E section was cut and the DCIS was graded by the study histopathologist (SEP) according to UK and College of American Pathologists guidelines [22]. Data on grade of DCIS were available from histopathology reports for 828 BCAC cases.

A subset of 81 ICICLE cases, graded in the pathology report and with a tumor block available, were examined to assess the reliability of the cytonuclear grade provided by the pathology reports. In the majority of cases (86.5 %) grade was concordant with the pathology report. Nine cases were re-graded as low/intermediate grade and two cases as high grade. As the study pathologist re-graded the samples on a single H&E section, rather than all the blocks from an individual case, and in some cases on reexcision specimens with residual disease rather than the original excision specimen, the grade reported in the pathology report, if available, was used for the purposes of this study.

ER status from local histopathology reports was available for 1,086 ICICLE cases. For the remaining 781 ICICLE cases where the tumor block was available, immunohistochemistry was performed on 3-µM sections, which were incubated at 60 °C for 1 h prior to automated staining using the VENTANA[®]. Estrogen receptor staining was carried out using CONFIRM[™] anti-estrogen receptor (SP1) rabbit monoclonal primary antibody (Catalog number 790-4324) with no variation to the recommended protocol. ER staining was scored by three independent reviewers (CP, VS, DLe) using the Allred method, and any discrepancies were reviewed by the study histopathologist (SEP). DCIS with an Allred score ≥ 3 was considered ER+ and DCIS with scores of 0-2 (approximately equivalent to <1 % of nuclei) was regarded as ER-. ER status was available on 965 cases from BCAC (Additional file 6).

Results

Assessment of known breast cancer susceptibility loci for association with DCIS

For the majority of known loci (n = 46) the risk allele for invasive breast cancer is the minor allele. For the ORs presented here the reference allele was set as the non-risk allele to make it clear whether the association with DCIS was in the same direction as previously published for invasive breast cancer. Thus, ORs for DCIS will be >1 if in the same direction as invasive disease and <1 if in the opposite direction.

Of the 76 known common breast cancer susceptibility loci genotyped on the iCOGS array, 51 were associated with DCIS (P < 0.05), with the effect in the same direction as previously reported in IDC (Fig. 1 and Additional file 7). Sixteen SNPs were significantly associated with DCIS (P < 0.00066) with three being genome-wide significant ($P < 5 \times 10^{-8}$, Table 1). The strongest associations were with for loci in *FGFR2* (rs2981579: OR 1.29, 95 % CI 1.24, 1.35; $P = 9.0 \times 10^{-30}$) and *TOX3* (rs3803662: OR 1.15, 95 % CI 1.1, 1.21; $P = 1.7 \times 10^{-8}$).

The case-only analysis (DCIS vs IDC) confirmed the shared genetic susceptibility between DCIS and IDC as none of the heterogeneity *P* values (*P*-Het) were significant after Bonferroni adjustment for 76 SNPs (Additional file 7). The case-only analysis (DCIS diagnosed at <50 years vs \geq 50 years of age) revealed one SNP (rs527616, 18q11.2) that was significantly associated with DCIS in younger women (*P*-Het_{<50/ \geq 50} = 0.0003) even though the overall *P* value for DCIS was not statistically significant after Bonferroni correction (OR 1.05, 95 % CI 1.01, 1.11; *P* = 0.020) (Additional file 8).

Assessment of known breast cancer susceptibility loci for association with DCIS by ER status

Following immunohistochemistry for ER in the ICICLE study samples, 1,484 cases (54 %) were classified as ER+ and 383 (14 %) as ER-. The ER data on BCAC DCIS were less complete with 664 (28 %) ER+, 301 (13 %) ER- and 1,387 cases (59 %) of unknown ER status (Additional file 6). Analysis by ER status confirmed that loci associated with ER+ IDC were also associated with ER+ DCIS (Fig. 2 and Additional file 9). These similarities were less clear for ER- DCIS and ER- IDC but this may be due to small numbers of ER- DCIS cases. A case-only analysis of ER+ vs ER- DCIS was not performed due to the small numbers of ER- cases.

SNP	Locus	
rs29815		
rs298158		
rs11814 rs37573		
rs10995		
rs38219		
rs380360		
rs55421		
rs29435 rs10771		
rs889312		
rs10941		
rs48498		
rs497370		_
rs480880		
rs75915 rs682852		
rs135374		
rs11780		
rs20462		
rs258880		
rs133870	42 2q35	_ →
rs647290		
rs424573		
rs600193		
rs178174		
rs12920 rs676264		
rs100690		
rs104720		
rs969344		
rs12422		
rs16857		
rs10119		
rs173569		
rs865686 rs72047		
rs61648		
rs941764	CCDC88C	
rs13329		_
rs282309		
rs99973		
rs223600		
rs704010 rs132810		
rs12710		
rs527610		-
rs112420		
rs14326	9 EBF1	*
rs79045		*
rs20163		
rs11820		
rs273610 rs11199		
rs66789		
rs65049		
rs143690		
rs155062		_
rs20424	RANBP1	*
rs23639		+
rs11075		
rs107593		
rs381719 rs17529		
rs12022		
rs39030		
rs376098		
rs97905		_
rs8170	MERIT40	
rs11977		
rs10454		_
rs238020		- <u>+</u> -
rs70727		
rs124930 rs132390		
rs112494		
1311240	55 ipi1.2	-
		.8 0.9 1 1.1 1.2 1.3 1.4

indicate that the association is in the same direction as previously published for invasive breast cancer

Table 1 Loci showing a significant association with ductal carcinoma in situ (DCIS) at P < 0.00066

Chromosome SNP Locus RAF		RAF	DCIS vs controls (meta-analysis)			IDC vs controls			Case-only DCIS vs IDC	
			Controls	OR	(95 % CI)	Р	OR	(95 % CI)	Р	<i>P</i> -Het
10	rs2981579	FGFR2	0.40	1.29	(1.24, 1.35)	9.0×10^{-30}	1.24	(1.21, 1.28)	6.1 × 10 ⁻⁶⁶	0.14
10	rs2981582	FGFR2	0.38	1.28	(1.23, 1.34)	1.8×10^{-27}	1.23	(1.20, 1.26)	2.1 × 10 ⁻⁵⁹	0.21
16	rs3803662	TOX3	0.26	1.15	(1.10, 1.21)	1.7 × 10 ⁻⁸	1.23	(1.20, 1.27)	1.5×10^{-50}	0.69
5	rs889312	MAP3K1	0.28	1.14	(1.09, 1.20)	6.9×10^{-8}	1.11	(1.08, 1.14)	2.2×10^{-14}	0.13
3	rs4973768	SLC4A7	0.47	1.13	(1.08, 1.18)	9.1 × 10 ⁻⁸	1.09	(1.07, 1.12)	8.2 × 10 ⁻¹³	0.58
5	rs10941679	5p12	0.25	1.14	(1.09, 1.20)	1.3×10^{-7}	1.14	(1.11, 1.18)	1.2×10^{-20}	0.90
3	rs3821902	ATXN7	0.13	1.16	(1.09, 1.23)	3.0×10^{-6}	1.06	(1.02, 1.09)	0.0030	0.33
19	rs4808801	SSBP4	0.65	1.12	(1.06, 1.18)	3.1 × 10 ⁻⁶	1.09	(1.05, 1.11)	3.5×10^{-9}	0.16
10	rs10995190	ZNF365	0.85	1.16	(1.09, 1.23)	4.1×10^{-6}	1.15	(1.11, 1.19)	7.5×10^{-16}	0.61
2	rs13387042	2q35	0.51	1.10	(1.05, 1.15)	1.1×10^{-5}	1.14	(1.11, 1.16)	8.3×10^{-25}	0.34
6	rs3757318	ESR1	0.07	1.20	(1.10, 1.30)	1.4×10^{-5}	1.16	(1.10, 1.21)	1.2 × 10 ⁻⁹	0.85
11	rs554219	CCND1	0.12	1.15	(1.08, 1.22)	2.8×10^{-5}	1.27	(1.22, 1.32)	6.4×10^{-38}	0.88
6	rs2046210	ESR1	0.34	1.10	(1.05, 1.15)	8.6×10^{-5}	1.09	(1.06, 1.12)	4.0×10^{-10}	0.32
12	rs10771399	PTHLH	0.88	1.15	(1.06, 1.23)	0.00021	1.18	(1.12, 1.22)	1.2×10^{-14}	0.53
8	rs11780156	8q24.21	0.16	1.11	(1.05, 1.18)	0.00027	1.10	(1.06, 1.14)	2.3 × 10 ⁻⁸	0.88
16	rs17817449	FTO	0.60	1.09	(1.03, 1.14)	0.00052	1.06	(1.04, 1.10)	5.9×10^{-7}	0.32

SNP single nucleotide polymorphism, IDC invasive ductal carcinoma, OR odds ratio; P-Het P value for heterogeneity; RAF risk allele frequency

Assessment of known breast cancer susceptibility loci for association with DCIS by grade

Grade data were available for 95 % of ICICLE DCIS cases; 1,635 (60 %) were of high cytonuclear grade and 943 (35 %) of low/intermediate grade. The grade data on the BCAC DCIS were less complete with data only available for 35 % of cases: 306 (13 %) high grade and 522 (22 %) low/intermediate grade cases (Additional file 6). Case–control analysis was performed separately on the low/intermediate and high grade subsets and a case-only analysis of low/intermediate grade vs high grade DCIS was performed to assess whether any of these loci were grade-specific.

Analysis of DCIS by grade revealed that although the majority of SNPs predispose to all grades of DCIS, some are grade-specific (Additional files 10 and 11). The two SNPs close to *CCND1* were strongly associated with low/intermediate grade DCIS (rs75915166, OR 1.36, 95 % CI 1.17, 1.59; $P = 7.2 \times 10^{-5}$; rs554219, OR 1.32, 95 % CI 1.18, 1.48; $P = 8.2 \times 10^{-7}$) and there was no association with high grade DCIS (Table 2). Case-only analysis confirmed that these loci were low/intermediate grade-specific (rs75915166, *P*-Het_{low/highgrade} = 0.00014; rs554219, *P*-Het_{low/highgrade} = 0.00013) and this was independent of ER status (adjusted for ER status rs75915166, P = 0.0050; rs554219, P = 0.019).

A similar-case-only analysis of IDC by grade confirmed that the two SNPs on 11q13.3 close to *CCND1* were also invasive grade 1/2-specific in IDC (rs75915166, OR 1.42, $P = 1.7 \times 10^{-30}$, P-Het = 2.8×10^{-10} ; rs554219, OR 1.39, $P = 4.7 \times 10^{-49}$, P-Het = 1.3×10^{-17}) and again were independent of ER status ($P = 1.3 \times 10^{-6}$, $P = 1.6 \times 10^{-6}$, respectively) (Additional file 12). In addition, other grade-specific loci were identified including three (rs2363956, rs8170 and rs10069690) specific to grade 3 invasive disease (Additional file 13).

rs10941679, 5p12 were borderline associated with low/intermediate grade DCIS (OR 1.26, $P = 2.1 \times 10^{-7}$, *P*-Het_{low/highgrade} = 0.0033). This locus has previously been shown to be associated with low grade progesterone receptor (PR) + IDC [23]. There was no evidence of any high grade DCIS specific loci (Additional file 11).

Search for new DCIS predisposition loci

All SNPs that were genome-wide significant ($P < 5 \times 10^{-8}$) in the meta-analysis were correlated with one of the known breast cancer predisposition loci. There were three SNPs that were not correlated with known loci at $P < 6 \times$ 10^{-6} (Table 3), all with very little evidence of an association with IDC.

Of these novel SNPs, rs12631593, 3p14.2, (an intronic variant in *FHIT*, chr3: 60726844) was the most strongly associated with DCIS (OR 1.21, 95 % CI 1.13, 1.29; $P = 5.5 \times 10^{-8}$). This SNP showed little association with IDC (OR 1.01, 95 % CI 0.97, 1.05; P = 0.54) and this was supported by the case-only analysis (*P*-Het_{DCIS/IDC} = 0.0048).



The other loci were on 22q13.2, rs73179023 (DCIS only: OR 0.85, 95 % CI 0.79, 0.90; $P = 1.1 \times 10^{-6}$; IDC only: OR 0.97, 95 % CI 0.93, 1.00; P = 0.060, P-Het_{DCIS/IDC} = 0.0099) and 7q21.3, rs13236351 (DCIS only: OR 1.30, 95 % CI 1.16, 1.46; $P = 5.7 \times 10^{-6}$; IDC only: OR 1.05, 95 % CI 0.99, 1.13; P = 0.13, P-Het_{DCIS/IDC} = 0.17).

These SNPs were genotyped in a validation study including a further 653 DCIS cases and 1,882 controls, however, for all three loci there was no evidence of an association (for rs12631593, rs13236351, and rs73179023, P = 0.49, 0.61, and 0.57, respectively) and none were genome wide significant following a meta-analysis of all data ($P = 7.8 \times 10^{-7}$, 2.9×10^{-5} , and 1.7×10^{-6} respectively) (Table 3).

Discussion

This study provides the strongest evidence to date for a shared genetic susceptibility between DCIS and IDC, based on 5,067 cases with pure DCIS (no invasive disease)

and 24,670 cases with IDC. It differs from previous BCAC analyses of DCIS, as it has included an additional 3,078 DCIS cases, excluded all cases of pure LCIS and has also compared DCIS to IDC rather than all invasive disease.

An important finding of this study is the lack of DCIS/ IDC-specific loci among the known breast cancer predisposition loci. Of the five breast cancer predisposition alleles originally reported by Easton et al. [24], three were shown to be associated with *in situ* (998 cases of DCIS and LCIS) disease (rs2981582-*FGFR2*, rs3803662-*TOX3*, rs889312-*MAP3K1*) with rs889312 showing a stronger association with DCIS (*P*-trend 0.007, per allele OR 1.30 for DCIS, per allele OR 1.13 for invasive disease). However, this finding of potential DCIS-specific loci was not confirmed in the Million women study which found no differential association with DCIS vs IDC for twelve breast cancer susceptibility loci, including rs889312, although their sample size was smaller (873 DCIS and 4,959 IDC) [12]. In the recent BCAC

Table 2 Association between rs75915166 or rs554219 and grade in ductal carcinoma in situ

	Meta-analysis				
	OR (95 % CI)	Р	Low/intermediate grade, number	High grade, number	Controls, number
rs75915166					
Low/intermediate grade vs controls	1.36 (1.17, 1.59)	7.2 × 10 ⁻⁵	1,465		35,521
High grade vs controls	0.92 (0.79, 1.08)	0.31		1,941	32,202
Case-only high vs low/intermediate grade					
Unadjusted	0.68 (0.55, 0.83)	1.4×10^{-4}	1,307	1,941	
unadjusted (only cases with ER status)	0.65 (0.51, 0.84)	1.1 × 10 ⁻³	791	1,360	
adjusted for ER status	0.68 (0.52, 0.89)	0.0050	791	1,360	
ER+ only	0.68 (0.55, 0.84)	5×10^{-4}	709	985	
rs554219					
Low/intermediate grade vs controls	1.32 (1.18, 1.48)	8.2 × 10 ⁻⁷	1,465		35,521
High grade vs controls	1.02 (0.91, 1.14)	0.75		1,941	32,202
Case-only high vs low/intermediate grade					
Unadjusted	0.75 (0.65, 0.87)	1.3×10^{-4}	1,307	1,941	
unadjusted (only cases with ER status)	0.75 (0.63, 0.88)	2.1 × 10 ⁻⁴	791	1,360	
adjusted for ER status	0.80 (0.67, 0.96)	0.019	792	1,360	
ER+ only	0.76 (0.65, 0.89)	6.7 × 10 ⁻⁴	709	985	

OR odds ratio, ER estrogen receptor

COGS analysis all 41 novel SNPs identified on the iCOGS chip had comparable ORs for invasive and in situ disease (based on data from 2,335 in situ, and 42,118 invasive cases), with the exceptions of rs12493607 (TGFBR2), and rs3903072 (11q13.1), for which associations seemed to be restricted to invasive disease [19]; however, we found no evidence of an IDC-specific association with these loci after correcting for multiple testing. A recent study investigating the association between 39 of the known breast cancer predisposition loci and breast cancer in situ (BCIS) suggested that rs1011970 (9p21.3, CDKN2BAS) had a stronger association with BCIS than invasive breast cancer (BC), P-Het_{BCIS/BC} = 0.0065. This trend remained in a DCIS vs BC analysis (P-Het_{DCIS/BC} = 0.021) [25]. Our data, however, do not support this finding (DCIS OR 1.08, 95 % CI 1.02, 1.14; *P* = 0.011; IDC OR 1.05, 95 % CI 1.0, 1.09; P = 0.0025, P-Het_{DCIS/IDC} = 0.33).

We have also shown for the first time that seven of the known invasive breast cancer predisposition loci not previously shown to be associated with DCIS have comparable ORs for IDC and DCIS: rs4973768 (*SLC4A7*), rs3821902 (*ATXN7*) [26], rs109 95190 (*ZNF365*), rs554219 (*CCND1*), rs3757318 and rs2046210 (*ESR1*).

This lack of DCIS/IDC-specific loci is in contrast to our previous study of lobular cancer in which we showed that there are loci that are specific to invasive lobular cancer (ILC), showing no association with lobular carcinoma in situ (LCIS) and there was also a suggestion of LCISspecific loci [16]. When we compare the DCIS data presented here to our previous LCIS analyses it reveals that there is some overlap between loci that are associated with ER+ DCIS and LCIS (Fig. 3 and Additional file 14). However, there are also some differences: rs6678914, *LGR6* and rs8656866, 9q31.2 are strongly associated with LCIS but there is little evidence of association with ER+ DCIS (*P*-Het_{DCIS/LCIS} = 7.4×10^{-5} and 6.6×10^{-4} , respectively). We have also previously shown that rs11249433, 1p11.2 and rs11977670, 7q34 have a stronger association with invasive lobular cancer than IDC [16]. These loci were only weakly associated with LCIS and were not associated with ER+ DCIS in this analysis.

Most association studies of invasive breast cancer involve subgroup analyses based on ER status. In contrast to invasive breast cancer, ER status in DCIS is not routinely assessed in all centers despite evidence from the NSABP B-24 trial of benefit from endocrine therapy in ER+ DCIS [7]. A national audit of DCIS in the UK revealed that ER status was assessed in only 50 % of DCIS cases and ER positivity in low and intermediate grade DCIS (P < 0.001) (ER+ high grade 69 %, intermediate grade 94 %, low grade 99 %) [27]. In order to overcome this issue we performed ER immunohistochemistry on the samples from ICICLE for which ER status was unknown. However, there was still a large amount of

Table 3 Potential new ductal carcinoma in situ susceptibility loci

Single nucleotide polymorphism	rs12631593	rs13236351	rs73179023
Chromosome	3	7	22
Position	60701884	97772513	43424477
Locus	FHIT	LMTK2	PACSIN2:TTLL1
Minor allele frequency	0.11	0.032	0.13
ICICLE DCIS phase I			
Odds ratio (95 % CI)	1.15 (1.04, 1.28)	1.31 (1.10, 1.56)	0.83 (0.75, 0.91)
Р	0.0088	0.0029	0.00020
BCAC DCIS			
Odds ratio (95 % CI)	1.25 (1.14, 1.36)	1.3 (1.12, 1.51)	0.86 (0.79, 0.94)
Р	1.0×10^{-6}	0.00060	0.0012
Meta-analysis phase I			
Odds ratio (95 % CI)	1.21 (1.13, 1.29)	1.3 (1.16, 1.46)	0.85 (0.79, 0.90)
Ρ	5.5×10^{-8}	5.7×10^{-6}	1.1×10^{-6}
Phase II DCIS			
Odds ratio (95 % CI)	0.93 (0.76, 1.14)	0.91 (0.63, 1.31)	0.95 (0.78, 1.15)
Ρ	0.49	0.61	0.57
Meta-analysis phase II			
Odds ratio (95 % CI)	1.18 (1.10, 1.25)	1.26 (1.13, 1.41)	0.86 (0.80, 0.91)
Р	7.8×10^{-7}	2.9×10^{-5}	1.7×10^{-6}
BCAC IDC			
Odds ratio (95 % Cl)	1.01 (0.97, 1.05)	1.05 (0.99, 1.13)	0.97 (0.93, 1.00)
Р	0.54	0.13	0.060
Case-only			
DCIS vs IDC <i>P</i> -Het	0.0048	0.17	0.0099

DCIS ductal carcinoma in situ, IDC invasive ductal carcinoma, BCAC Breast Cancer Association Consortium, ICICLE Study to investigate the genetics of in situ carcinoma of the ductal subtype, P-Het P value for heterogeneity

missing data on ER status in the BCAC cases, resulting in only 684 ER– DCIS cases being available for analysis, making it difficult to draw definitive conclusions about ER– DCIS. In essence the findings are similar to invasive breast cancer, with ER– and ER+ DCIS having different genetic susceptibility profiles and ER+ DCIS having a very similar profile to ER+ IDC.

Cytonuclear grade of DCIS is used by many clinicians to select those cases most likely to benefit from radiotherapy despite the fact that grade has not been shown to be a good predictor of recurrence. In the UK audit of DCIS, grade data were available for 99 % of DCIS cases, with 59 % classified as high grade, 29 % as intermediate and 11 % as low grade [27]. Similarly, in our study data on grade were available for 95 % of cases in ICICLE. In invasive disease only a minority of predisposition loci have been shown to be grade specific; rs2981582 (*FGFR2*) and rs13281615 (8q24) [28, 29] and rs10941679 (5p12) [23]. We have shown that analysis of DCIS by grade reveals other known loci that are grade specific. The loci with the strongest association with grade were SNPs on 11q13, which had a stronger association with low/intermediate grade DCIS and IDC than high grade lesions. The finding of a strong association with low and intermediate grade ductal carcinomas that is independent of ER status in both DCIS and IDC for these loci is novel. rs614367 was the first locus on 11q13 shown to be associated with invasive breast cancer [30]. Fine mapping of the region subsequently identified two independent signals (rs554219 and rs78540526, $r^2 = 0.38$), which are the loci reported in this analysis. Functional analyses demonstrated that the risk variants modify enhancer and silencer elements, with the likely target gene being *CCND1* [31].

A study of 150 cases of subsequent breast cancer (invasive and in situ) after DCIS observed significant association for both grade and ER status between the index DCIS and the subsequent breast cancer (whether ipsilateral or contralateral), suggesting that women with DCIS are at risk of developing subsequent breast cancers



of a similar phenotype [32]. This finding supports the genetic predisposition data presented here, with ER and grade-specific loci in DCIS having similar specificity in IDC.

Although we did not identify any novel loci that reached genome wide significance, we did identify three potential novel DCIS predisposition loci, two of which were DCIS-specific (rs12631593, rs73179023), and therefore need further investigation in other cohorts of DCIS. As at least 45 % of patients with IDC have associated DCIS present at diagnosis consistent with direct precursor behavior, it may seem biologically implausible that an SNP predisposes to DCIS but is not associated with IDC. However, it is possible that there is a subset of patients with DCIS with very low probability of progression. If the finding of DCIS-specific predisposition loci were confirmed in other studies, identifying such a subset of patients with low-risk DCIS would be clinically valuable.

Conclusion

In conclusion this is the largest study to assess genetic predisposition in DCIS and shows that the majority of invasive breast cancer predisposition loci also predispose to DCIS. It highlights that, as for invasive disease, different SNPs predispose to ER+ and ER- DCIS. In addition it shows the importance of grade in both DCIS and IDC.

Additional files

Additional file 1: Study information for the Breast Cancer Association Consortium (*BCAC*) participating studies. (DOCX 28 kb) Additional file 2: Sample information for the SEARCH, UKBGS, SBCS, and BBCS studies. (DOCX 15 kb)

Additional file 3: Number of studies and individuals included in analyses of ductal carcinoma *in situ (DCIS)* and invasive ductal carcinoma *(IDC)*. BCAC Breast Cancer Association Consortium. (XLSX 12 kb)

Additional file 4: Principal component analysis (PCA) results from the study to investigate the genetics of *in situ* carcinoma of the ductal subtype (*ICICLE*). (PPTX 142 kb) Additional file 5: Quantile-quantile plots from the study to investigate the genetics of in situ carcinoma of the ductal subtype (*ICICLE*). *SNP* single nucleotide polymorphism. (PPTX 125 kb)

Additional file 6: Grade, estrogen receptor (*ER*) status, and age groups in patients with ductal carcinoma in situ (*DCIS*). *BCAC* Breast Cancer Association Consortium, *ICICLE* study to investigate the genetics of *in situ* carcinoma of the ductal subtype. (DOCX 16 kb)

Additional file 7: Association between ductal carcinoma in situ (DCIS) and known breast cancer predisposition loci. *IDC* invasive ductal carcinoma, *P-Het P* value for heterogeneity, *SNP* single nucleotide polymorphism, *OR* odds ratio. (XLSX 19 kb)

Additional file 8: Age-specific case-only analysis of patients with ductal carcinoma *in situ* (DCIS) diagnosed at age <50 vs \geq 50 years. *P-Het P* value for heterogeneity, *SNP* single nucleotide polymorphism, *OR* odds ratio. (XLSX 17 kb)

Additional file 9: Associations between the known breast cancer predisposition loci and estrogen receptor-positive (*ER+*) or *ER–* ductal carcinoma *in situ* (*DCIS*). *P-Het P* value for heterogeneity, *SNP* single nucleotide polymorphism, *OR* odds ratio. (XLSX 19 kb)

Additional file 10: a, b Known breast cancer predisposition loci for low/intermediate grade (*black*) and high grade ductal carcinoma in *situ* (DCIS) (*gray*). Due to the large number of single nucleotide polymorphisms (*SNPs*), the plot is split for better visual representation into two different sections (a and b) with a descending order of effect size for the low/intermediate group. *OR* odds ratio. (ZIP 20 kb)

Additional file 11: Associations of the known breast cancer predisposition loci for high and low-intermediate grade ductal carcinoma *in situ (DCIS). P-Het P* value for heterogeneity, *OR* odds ratio. (XLSX 18 kb)

Additional file 12: Association of rs75915166 and rs554219 with grade in invasive ductal carcinoma (*IDC*). *P-Het P* value for heterogeneity, *OR* odds ratio. (XLSX 9 kb)

Additional file 13: Associations between the known and novel breast cancer predisposition loci and invasive ductal cancer, by estrogen receptor (*ER*) status and grade. *OR* odds ratio. (XLSX 24 kb)

Additional file 14: Association between the known breast cancer predisposition loci and estrogen receptor-positive (*ER+*) ductal carcinoma *in situ* (*DCIS*) or lobular carcinoma *in situ* (*LCIS*). *P*-Het *P* value for heterogeneity, *SNP* single nucleotide polymorphism, *OR* odds ratio. (XLSX 19 kb)

Abbreviations

ABCS: Amsterdam Breast Cancer Study; BBBC: Bavarian Breast Cancer Cases and Controls; BBCS: British Breast Cancer Study; BC: breast cancer; BCAC: Breast Cancer Association Consortium; BCIS: breast carcinoma in situ; CI: confidence interval; COGS: Collaborative Oncological Gene-Environment Study; DCIS: ductal carcinoma in situ; ER: Estrogen receptor; H&E: hematoxylin and eosin; HWE: Hardy Weinberg equilibrium; ICICLE: study to investigate the genetics of in situ carcinoma of the ductal subtype; IDC: invasive ductal carcinoma; LCIS: lobular carcinoma in situ; MAF: minor allele frequency; OR: odds ratio; PCA: principal component analysis; *P-Het: P* value for heterogeneity; SNP: single nucleotide polymorphism.

Competing interests

The authors declare that there are no conflicts of interest.

Authors' contributions

The study was conceived by ES and RR. Analysis and genotyping in ICICLE was performed by ES. Meta-analyses were performed by MGC. The manuscript was prepared by ES. EJS, RR and IT conceived and designed the experiments. CP, VS, DLe, EP, AGN, DCT, DV, FB, JD, and AMD performed the experiments. CP, MNB, MKB, QW, KM, IT, MGC, and EJS analyzed the data. CP, MNB, VS, KK, PGo, MC, DLe, EP, NO, AC, SSC, IdSS, JP, AS, MJS, MKB, QW, JD, KM, JB, AGN, DCT, DV, JLi, JF, VK, ALBD, PS, JS, RLM, GGG, SM, AL, TB, HBra, MCS, JLH, TD, NVB, MK, UH, RKS, AMe, HBre, VA, RW, KP, PAF, MVB, JLu, AJ, AMM, ILA, RAEMT, PD, LLM, CAH, AMA, VMK, PR, PP, FM, BB, CHMvD, AH, NM,

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Authors' information

Study was conceived by ES & RR, analysis & genotyping of ICICLE performed by ES, meta-analyses performed by MGC, manuscript prepared by ES.

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BBCC	Friedrich-Alexander-Universitat Erlangen-Nurnberg Medizinische Fakultat Ethik-Commission
BBCS	South East Multi-Centre Research Ethics Committee
BIGGS	Galway University College Hospital Clinical Research Ethical Committee
BSUCH	Medizinische Fakultat Heidelberg Ethikkommission
CECILE	Comite Consultatif de Protection des Personnes dans la Recherche Biomedicale de Bicetre
CGPS	Kobenhavns Amt den Videnskabsetiske Komite
CNIO-BCS	Hospital Universitario La Paz Comite Etico de Investigacion Clinica
ESTHER	Ruprecht-Karls-Universitat Medizinische Fakultat Heidelberg Ethikkommission
GC-HBOC	Ethik-Kommission der Medizinischen Fakultat der Universitat zu Koln
HEBCS	Helsingin ja uudenmaan sairaanhoitopiiri (Helsinki University Central Hospital Ethics Committee)
HMBCS	Medizinische Hochschule Hannover Ethik-Kommission
ICICLE	Southampton and South West Hampshire Research Ethics Committee A (MREC 08/H0502/4)
КВСР	Pohjois-Savon Sairraanhoitopiirin Kuntayhtyma Tutkimuseettinen Toimikunta
kConFab/AOCS	kConFab: The Queenland Institute of Medical Research Human Research Ethics Committee (QIMR-HREC)
	AOCS: Peter MacCallum Cancer Centre Ethics Committee
MARIE	Ruprecht-Karls-Universitat Medizinische Fakultat Heidelberg Ethikkommission
MBCSG	Comitato Etico Indipendente della Fondazione IRCCS "Istituto Nazionale dei Tumori"
MCBCS	Mayo Clinic IRB
MEC	University of Southern California Health Sciences Campus IRB
OBCS	Ethical Committee of the Medical Faculty of University of Oulu and Northern Ostrobothnia Hospital District Ethical Committee
OFBCR	Mount Sinai Hospital Research Ethics Board
ORIGO	Medical Ethical Committee and Board of Directors of the Leiden University Medical Center (LUMC)
pKARMA	Regionala Etikprovningsnamnden i Stockholm (Regional Ethical Review Board in Stockholm)
RBCS	Medische Ethische Toetsings Commissie Erasmus Medisch Centrum
SBCS	South Sheffield Research Ethics Committee
SEARCH	Multi Centre Research Ethics Committee (MREC)
SZBCS	Komisji Bioetycznej Pomorskiej Akademii Medycznej
UKBGS	South East Multi-Centre Research Ethics Committee

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