COMMENTARY

Epidemiology, Biology, Treatment, and Prevention of Ductal Carcinoma In Situ (DCIS)

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Approximately 50,000 cases of ductal carcinoma in situ (DCIS) are diagnosed in the United States each year (1). The term “DCIS” encompasses a highly heterogeneous group of lesions that differ in their clinical presentation, histologic and biologic features, and outcome. Although DCIS is considered to be a precursor to invasive breast cancer, 14–53% of DCIS will not progress to invasive breast cancer, and thus it is considered a nonobligate precursor (2–6). As a result, there remains considerable uncertainty about optimal clinical management at the individual patient level.

In May 2017, the Dana-Farber/Brigham Cancer Center brought together clinicians and researchers from multiple institutions to discuss the current trends and identify and address the key questions (Supplemental Table 1, available online) that must be answered to improve our understanding and clinical management of DCIS.

Part I: Current Trends, Outcomes, Detection, and Pathology

Treatment Outcomes and Trends

Surgery is the standard of care for DCIS; however, until recently there were limited data examining the impact of surgery, with or without radiation, on survival by grade or size of DCIS. Data were also limited on the use of adjuvant therapies for DCIS nationally and whether these therapies truly improve outcomes. To address this gap, Sagara and colleagues conducted a study to determine the survival benefit of surgical treatment by nuclear grade in patients with DCIS (7). Using data from the Surveillance, Epidemiology, and End Results (SEER) database, they analyzed a cohort of women diagnosed with DCIS by biopsy only. Overall, the study showed no breast cancer-specific
survival benefit of surgery for women with low-grade disease; however, intermediate- or high-grade DCIS patients did experience a survival benefit from surgery.

The Early Breast Cancer Trialists’ Collaborative Group examined the efficacy of radiotherapy (RT) following breast-conserving surgery (BCS) for DCIS in a meta-analysis of four randomized controlled trials (8), and reported that RT reduced the rate of local recurrence by ~50%, irrespective of tumor type and patient characteristics. However, RT did not improve overall survival or breast cancer-specific survival. In contrast, Sagara et al used SEER data along with patient prognostic score to investigate the benefit of RT among patients treated between 1988 and 2007 (9). Propensity score analysis of the overall cohort demonstrated a small yet statistically significant survival benefit (0.3%) associated with the receipt of radiation (9), reflecting the power of a large population-based analysis and suggesting that RT may modestly improve survival for some patients with DCIS.

Using the American College of Surgeons’ National Cancer Database, Sagara et al also investigated factors associated with the use of adjuvant RT and/or endocrine therapy (ET) following BCS in women with DCIS (10). This study demonstrated that both clinico-pathologic and demographic factors influence the use of adjuvant therapy. For hormone receptor (HR)-positive DCIS treated with BCS, there has been a shift towards decreasing use of RT and increasing use of ET alone or ET in combination with RT after BCS. In contrast, for HR-negative DCIS, the proportion of women receiving RT after BCS has increased over the last decade. Further, in a low-risk cohort—defined as patients older than 60 years of age with low-grade HR-positive DCIS lesions, less than 16 mm in size, excised to negative margins—the use of RT and/or ET with RT has decreased (Figure 1). These national trends suggest that there is a DCIS patient population and their physicians who are receptive to de-escalating therapy.

**Controversies in Imaging in DCIS**

Prior to 1985, DCIS represented 2% of all breast cancer. When mammography screening began, incidence increased and, based on SEER data from 2007–2013, one-third of breast cancers detected by screening mammography in current practice are DCIS. Some see this increased detection as a problem, leading to overtreatment for DCIS patients, and call for reduced screening mammography (11). In recent years, breast imaging has shifted from 2D mammography to 3D mammography and magnetic resonance imaging (MRI). On 2D mammography, breast imagers were looking for patterns suggestive of cancer, and a focus was on calcifications. With the introduction of 3D mammography, or tomosynthesis, the overall cancer detection rate has not varied dramatically, but there has been a lower percentage of DCIS diagnoses and a higher percentage of invasive cancers detected compared with 2D (12–14). This suggests that tomosynthesis may be less sensitive for detecting DCIS and/or that mammographers are now focusing more attention on identifying imaging features that are more likely to be associated with invasive cancers and possibly less attention to microcalcifications. Despite this, we have not seen a significant decrease in interval cancers with tomosynthesis, suggesting that this shift in detection patterns may not be leading to improved clinical outcomes.

Several studies of mammography, MRI, and ultrasound confirm that MRI has the highest sensitivity for detecting DCIS (15,16). Kuhl et al. calculated the performance of imaging modalities based on DCIS grades (16). When compared with mammography, MRI was superior in detecting both low- and high-grade DCIS, but MRI sensitivity was especially evident in high-grade DCIS (low-grade ~ 80% MRI detection/60% mammography detection vs high-grade ~ 98%/52%). However, the type of mammography used in this study for three of the five years was film-screen, known to have inferior sensitivity for DCIS detection to full-field digital mammography, which is now in widespread use.

The ECOG-ACRIN DCIS 4112 study aims to identify women with DCIS who may be managed less aggressively without sacrificing excellent outcomes. This study investigates the use of preoperative breast MRI and its impact on surgical treatment decisions. Early findings suggest that conversion to mastectomy was common, with nearly one in five patients ultimately undergoing mastectomy, and that breast MRI findings accounted for less than one-half of these conversions (Table 1). Patient preference is a strong factor for converting to mastectomy for a wide range of reasons. For patients who remained candidates for wide local excision after MRI, 96.4% achieve successful wide local excision as the final surgical procedure. MRI also highlights other lesions, of which more than one-half are benign (17). These false positive findings require careful management with effective communication between the radiologist, surgeon, and patient to reduce anxiety and avoid unnecessary mastectomies.

It is also important to note that modern imaging has the potential not only to identify lesions that may be cancer, but also to evaluate information beyond the lesions. For example, features of the breast tissue around the DCIS may be potential biomarkers for risk of recurrence or invasive disease. Future research must determine how and in what context we can use this type of information clinically.

**Risk Prediction and Uncertainty**

After decades of research, it is still not possible to reproducibly identify which DCIS lesions will progress to invasive disease and which are unlikely to progress and, correspondingly, which patients can be managed safely with excision alone or no treatment beyond the diagnostic biopsy. Young age has been shown to be a risk factor for local recurrence in patients with DCIS. There appear to be no differences in distribution of pathology-related factors (grade or necrosis) according to age, though younger patients are more likely to have greater extent of disease and present with a palpable mass lesion (18). Additionally, women with symptomatic presentation (ie, palpable mass in breast) are more likely to progress to invasive breast cancer (19).

The association between specific genetic changes and grade could provide insights into the biology of DCIS that influences pathologic classification and clinical management. All of the intrinsic molecular subtypes identified in invasive cancers are also seen in DCIS (20). However, among patients with DCIS, a slightly greater proportion are classified as HER2 enriched. Two groups have investigated whether certain DCIS phenotypes are associated with an increased risk of invasive cancer. Williams et al. (21) concluded that all phenotypes were associated with an increased risk compared with the luminal A phenotype, and the Cancer Research Network showed only the HER2-enriched group was at increased risk (Laurel A. Habel, Ninah Achacoso, Stuart J. Schnitt, Laura C. Collins, Monica Morrow, Reina Haque, Larissa Nekhlyudov, Suzanne W. Fletcher, Allen M. Gown, Lynn Goldstein, Charles P. Quesenberry, Jr., unpublished data). Many studies have shown a relationship between various histologic features of DCIS and clinical outcome following BCS; however, the findings are often conflicting and thus the relative importance of the various histologic features is still not well defined.
Risk stratification based on pathologic factors remains elusive and this confounds communication of risk to patients. The USC/Van Nuys Prognostic index combines tumor size, grade, and margin status with patient age to estimate local recurrence risk and benefit from RT. Yet, this tool is challenging to apply in clinical practice due to the stringent sampling requirements, making it useful in only a minority of cases. Silverstein et al. argue that if the lesion is adequately excised (margin width > 10 mm), a patient’s risk of local recurrence is unaffected by nuclear grade, presence of comedo necrosis, lesion size, or addition of RT (24). However, given the surgical and cosmetic implications of such wide margin widths, this is difficult to achieve for all patients. A Memorial Sloan Kettering Cancer Center study shows that patients with a higher volume of disease near the margin derive a greater benefit from the addition of RT (25). Other data suggest patients with small, low-grade lesions may be adequately treated with wide excision only (26,27).

The Memorial Sloan Kettering Cancer Center nomogram is another tool that combines 10 clinico-pathologic factors to generate 5- and 10-year risks of ipsilateral breast cancer recurrence. This validated tool estimates outcomes with or without RT and/or ET. This score has a tendency to overpredict recurrence in patients treated with RT and underpredict recurrence in patients treated with BCS alone (28–30).

The Oncotype DCIS score is a commercial risk assessment test that is prognostic only. However, in two studies that have assessed the prognostic value of this score, patients in the low-risk group still have local recurrence rates over 10% at 10 years (31–33). More recent data suggest that integrating the DCIS score with tumor size and patient age helps further refine risk assessment. The DCISionRT test has also been recently developed to predict ipsilateral breast events after DCIS and benefit from RT (34). This test is currently undergoing further validation in two ongoing prospective clinical studies.

In summary, currently no histopathologic features of DCIS consistently provide accurate risk prediction for progression to invasive carcinoma. Several challenges were noted during the group discussion: that large databases with adequate outcomes including clinical and pathologic details have not been widely available, and DCIS outcomes are not the same today as in the older studies included in the prospective randomized trials. Several potential research directions were also discussed and are detailed in Supplemental Table 2 (available online).

**Part II: State of the Science**

**Biology of DCIS and the Role of the Microenvironment**

Little progress has been made over the last 20 years in reproducibly distinguishing biologically favorable from unfavorable DCIS. Given that few differences have been found at the genomic and transcriptomic levels between the cells of DCIS and invasive breast cancers of equivalent grade, it is possible that studying the DCIS microenvironment may be the key to understanding the biological progression from DCIS to invasive cancer.
By definition, the neoplastic cells in DCIS are present within the mammary ductal-lobular system and have not spread outside these sites. In the early stages of progression, immune cells may help to eliminate the cancer cells. In the equilibrium stage, the tumor is still somewhat controlled by the immune system; invasion/progression occurs when the tumor cells escape from the immune system. Understanding the mechanisms of escape in DCIS, which might be immune-mediated, may lead to more options for prevention.

Myoepithelial cells, which produce the basement membrane of the ducts and have a tumor suppressor function, are still present in DCIS (these cells are not present in invasive cancer) but the gene expression profiles and immunophenotype of myoepithelial cells in DCIS differ from myoepithelial cells in normal breast tissue (35). Myoepithelial cells prevent invasive progression both due to forming a structural barrier and also by expressing many anti-invasive and tumor suppressive genes (35–37). Therefore, myoepithelial cells can be viewed as “gatekeepers” of invasive progression, and alterations in myoepithelial cells seen in DCIS could predict risk of invasive recurrence (38). As many of the genes differentially expressed between DCIS-associated and normal myoepithelial cells encode for secreted proteins, it is possible that the myoepithelial cells orchestrate the microenvironmental changes, including changes in the immune microenvironment, that are present in DCIS compared with normal breast tissue.

When comparing normal breast tissue to DCIS and invasive cancers, the presence of myoepithelial cells, leukocytes, macrophages, cytotoxic T cells, and helper T cells in the microenvironment are different. Interestingly, Gil Del Alcazar and colleagues reported a decrease in activated cytotoxic T cells in invasive tumors compared with DCIS and at the same time also observed an increase in the expression of immune checkpoint proteins such as PD-L1 and CTLA4 with invasive progression (39). In the case of PD-L1 this included a selection for cancer cells that have amplification of the gene encoding for PD-L1 (39).

The microenvironment is diverse and this diversity shifts as the cancer progresses. There are measurable differences between these factors when comparing normal tissue, tissue from patients with DCIS and tissue from patients with invasive disease, as well as tissue from low-grade DCIS patients versus high-grade DCIS patients. If we can pinpoint the significance of the biologic changes in this environment, we may be able to distinguish a DCIS that will progress to invasive cancer from a DCIS that will not progress and guide patient management accordingly.

**Vaccines and Prevention**

Historically, vaccines were evaluated in patients with metastatic disease. Although early-phase trials showed that vaccination could generate antigen-specific immune responses, there was minimal evidence of clinically meaningful activity (40). Vaccinating patients with metastatic disease is a challenge due to extent of disease burden and the immunosuppressive microenvironment, which hampers T cell activity. To address these limitations, investigators have put forward the hypothesis that cancer vaccines may be more effective in a minimal disease setting to prevent disease recurrence after standard therapy or for primary prevention.

The goal of vaccinating in the adjuvant setting is to elicit a memory immune response that could be reactivated if cancer cells are detected in order to eliminate those cells before they can become established as recurrent disease. While a simple vaccination strategy may not be effective as secondary prevention, researchers have questioned whether it may be possible to administer a vaccine for primary prevention. This strategy is currently being used with the administration of the human papillomavirus vaccine to prevent human papillomavirus-associated malignancies. However, targeting tumor antigens rather than a virus is complex. Initially, it was thought that nonviral, nonmutated tumor antigens are too similar to self-antigens and targeting them may lead to autoimmunity. However, studies based largely on melanoma antigens have implied that autoimmunity is required for antitumor effect. In addition, epidemiologic data suggest that an immune response against epithelial antigens stimulated early in life either through the development of a childhood disease such as chicken pox or mumps, due to febrile illnesses, or via childhood vaccination, may decrease the risk of developing a malignancy (41–45). Therefore, it is possible that vaccines would be successful in preventing progression in patients with premalignant lesions (43).

Vaccination may be most effective in DCIS and atypical ductal hyperplasia before tumor cells are genetically unstable and rapidly dividing. Czerniecki and colleagues conducted a neoadjuvant study using a HER2-targeted dendritic cell vaccine administered to patients with a biopsy diagnosis of DCIS before surgery (46,47). Compared with pretreatment biopsies, surgical specimens showed an increase in CD4+ and CD8+ T cells after vaccination and a decrease in HER2 expression, suggesting the possibility of using vaccination to elicit an antigen-specific, anti-tumor immune response at the earliest stages of disease. Another trial, VADIS (NCT02636582), is a phase II trial investigating the E75þGM-CSF peptide vaccine administered three times before surgery in DCIS patients (48). The primary endpoint is the generation of E75-CTL T cells in vaccinated patients. Secondary endpoints include toxicity, epitope spreading, T cell functional capacity, and histologic response. Evaluating vaccines in patients with DCIS is an initial step toward developing a truly preventive breast cancer vaccine.

**Prevention of Invasive Breast Cancer: New Opportunities**

The main limitation of current prevention studies is that outcomes may be rare, and time to events can be prolonged; thus it may take many years for definitive conclusions to be drawn from these studies. Therefore, researchers are investigating surrogate outcomes (eg, changes in mammographic density, or molecular and immune biomarkers) for prevention. Investigators have conducted studies with agents that are unlikely to cause unwanted side effects, but these agents are also less likely to lead to large differences in risk (ie, dietary components such as flaxseed). By studying surrogate outcomes, the hope is to identify prevention agents that are tolerable and can be scaled.

Vitamin D, a steroid hormone (Alliance 70806) (49), and metformin, an antidiabetic drug (Alliance A221102) (50), are potentially promising agents for breast cancer prevention. Many women already take vitamin D for bone health and it is well tolerated; previous studies suggest the effectiveness of metformin to improve metabolic factors (51) and regulate levels of pAKT and pAMPK (52).

Given the results of the IBIS trial (53), which showed a long duration of benefit and significant breast cancer risk reduction for women who take tamoxifen, there is interest in finding new ways to use tamoxifen as a prevention agent. The Afimoxifene
in Reducing the Risk of Breast Cancer in Women With Mammographically Dense Breasts study (NCI) (54) is a randomized, phase II trial investigating the efficacy of tamoxifen topical gel applied to the breast area. Preliminary analysis reveals women were more willing to try transdermal tamoxifen than oral tamoxifen (55). In BRCA1 mutation carriers, there are ongoing prevention studies investigating bezalidoxime an an estrogen receptor degrader (56) and denosumab as a RANK ligand inhibitor (57).

Challenges identified during our group discussion included the heterogeneity in DCIS samples, which raises many issues related to the use of limited core biopsy samples and the need for fresh tissue to continue studying the underlying biology of progression of DCIS. Further, prevention trials have many challenges including historically low accrual, need for long-term treatment and follow-up, and well-studied but not lucrative drugs leading to high clinical trial costs. However, evaluation of these drugs as well as alternative approaches to prevention such as cryotherapy or intraductal injection of anti-tumor agents to elicit immediate and long-term immune rejection of lesions may yield important insights into DCIS prevention and treatment.

Part III: Risk Perceptions, Communication, and Decision-Making

Risk Perceptions, Communication, and Decision-Making for DCIS

The use of BCS as an alternative to mastectomy for DCIS gained acceptance in the 1990s (58). Randomized trials have demonstrated that adding RT after BCS reduces both the risk of having an invasive recurrence and a DCIS recurrence (59–62); however, a meta-analysis of these trials reveals that the use of RT does not improve survival (8). Differences in the interpretation of these data or values associated with local versus survival benefits have led to large regional variation in the use of RT for DCIS (Figure 2) (63,64).

Many attempts have been made to identify patients at low risk of local recurrence after BCS alone. However, there is little consensus on how best to combine classical clinico-pathologic characteristics in these analyses. As noted previously, the Oncotype DCIS score was developed to provide a gene expression assay for potentially improved characterization of risk of local recurrence, but how best to incorporate this assay in clinical practice remains uncertain. Raldow et al. sought to determine the cost-effectiveness of different treatment strategies using the Oncotype DCIS score (65). The authors used a Markov model to simulate 10-year outcomes for 60-year-old women eligible for the ECOG ES194 study (32) and determined the cost-effectiveness of using genomic testing after BCS for all women versus treating all women with excision alone or excision with RT (66). None of the treatment strategies employing the genomic test or strategies treating all women with RT was cost-effective relative to excision alone. Sensitivity analyses revealed that the most cost-effective strategy was highly sensitive to the utility of being without disease after excision alone or after excision and RT. This finding highlights the importance of engaging patient preferences in the treatment decision process.

Physicians note that decision-making about treatment for DCIS is very difficult for patients, who have a tendency to overestimate their risk of recurrence (67). In one study, more than 25% of DCIS patients believe that there is at least a moderate likelihood of DCIS spreading to other places in their body (68). These data underscore the need for patient education about treatment outcomes to improve the quality of decision-making about treatment. Decision-making is further complicated in that the outcome most important to an individual patient or stakeholder group may vary. Invasive breast cancer diagnosis is only one such outcome—other important outcomes include the likelihood of undergoing further breast biopsy or surgery, breast preservation, chemotherapy, and financial costs.

To help inform patients of the risks associated with different treatment options for DCIS, Punglia et al. have created a web-based decision aid: www.onlineDeCISion.org (69). Ongoing modifications to the tool will allow physicians to view outcomes by patient age and to tailor results by other treatment and risk factors if they are available (eg, age, grade, ER status) (70). One important message of the decision aid is that survival outcomes are essentially the same whichever treatment is chosen. With this information, patients may be able to better evaluate treatments based on their preferences and their tolerance for recurrence, versus the inconvenience or side effects of treatment, and improve the quality of their DCIS decisions.

Are We Ready for De-Escalation of Treatment for DCIS?

Ryser and colleagues conducted a usual care versus active surveillance analysis using SEER data on women with pure DCIS (Supplemental Figure 1, available online) (71). The main outcome measure was the probability of breast cancer death at 10 years. Under active surveillance, younger women had a higher risk of dying from breast cancer when compared with older women. However, by age 70 years, patients have much higher competing causes of mortality regardless of DCIS treatment choice. This does suggest that there may be a group of patients where active surveillance may be reasonable. Nevertheless, there is the concern of unrecognized invasive cancer, which has been reported in approximately 25% of patients diagnosed with DCIS on core needle biopsy (72). Therefore, patients can choose between the standard treatment options for DCIS (BCS, mastectomy, RT, and ET) or living with an increased risk of breast cancer-specific mortality ranging from 0.2% to 2.6% at 10 years. It is possible that some patients may consider this level of risk small and would prefer to forgo aggressive treatment, particularly in the face of preexisting comorbidities. Patient participation in evaluation of these trade-offs is crucial.

Currently there are three randomized controlled trials of active surveillance open to low-risk DCIS patients: LOW Risk DCIS (LORD), LOW Risk DCIS (LORIS) and Comparison of Operative to Monitoring and Endocrine Therapy (COMET) (Table 2) (73–77). In all three trials, patients are randomized between two treatment arms: 1) standard treatment regimen, or 2) no intervention, with close monitoring with mammography. The primary outcome for LORD and LORIS is ipsilateral invasive cancer-free survival; the primary endpoints for COMET are invasive cancer diagnosis, overall survival, disease-specific survival, quality of life, fear of cancer recurrence, and body image. Active surveillance is not encouraged outside of a clinical trial context, and this treatment strategy is not appropriate for patients with high-grade or extensive DCIS, palpable disease, mass on imaging, or other specific breast signs or symptoms.

The many challenges that remain also include how patient treatment decisions, and associated trial accrual, may be affected by the language used by health care professionals when
discussing DCIS and the difficulty in conveying uncertainty in outcome estimates to patients. Moreover, does randomization to active surveillance put a woman at risk for needing more extensive surgery later that may be more deforming than BCS? This is an important endpoint that will be collected in the active surveillance studies.

Table 2. Inclusion and exclusion criteria for the COMET, LORIS, and LORD trials (75)*

| CRITERIA                     | COMET                          | LORIS                         | LORD                          |
|-----------------------------|--------------------------------|-------------------------------|-------------------------------|
| Inclusion criteria          |                                |                               |                               |
| Age, y                      | ≥40                            | ≥46                           | ≥45                           |
| Nuclear grade               | Low and intermediate           | Low and intermediate          | Low                           |
| Morphology                  | Calcifications only            | Calcifications only           |                          |
| Hormone receptor status     | ER and/or PR positive, plus HER2 negative if performed | N/A                           | N/A                           |
| Exclusion criteria          |                                |                               |                               |
| History of cancer           | Exclude if invasive breast cancer | Exclude if invasive breast cancer or ipsilateral DCIS | Exclude if any cancer except in situ of the cervix or basal carcinoma of the skin |
| Symptomatic                 | Exclude                        | Exclude                       | Exclude                       |
| Comedonecrosis              | Exclude*                       | Exclude                       | N/A                           |
| Synchronous invasive cancer | Exclude                       | Include                       | Exclude                       |
| Bilateral DCIS at presentation | Include                     | Include                       | Exclude if family with BRCA 1/2 |
| High risk                   | Include                        | Exclude if high risk per NICE guidelines (76) | N/A                           |
| History of chemoprevention  | Exclude                        | N/A                           | N/A                           |

*Criteria deemed not applicable (N/A) are not mentioned in the inclusion or exclusion criteria of the study protocols. The table reports the data included in reference (75) in regards to the COMET trial exclusion criteria; however, the criteria were recently updated. Comedonecrosis no longer an exclusion criteria. The trial now allows any patients with low or intermediate grade DCIS. COMET – Comparing Operative to Medical Endocrine Therapy for low-risk DCIS; DCIS – ductal carcinoma in situ; LORD – Low Risk DCIS; LORIS – Low RISk DCIS; NICE – National Institute for Health and Care Excellence.
Future Research Directions
Patients with DCIS have excellent breast cancer-specific survival, irrespective of their choice for local therapy. Patients are not generally dying from the disease, so the question remains: how best do we treat each individual patient? We identified several take-aways from the retreat. First and foremost, patient preferences are paramount in treatment decisions regarding DCIS. We must strive to better educate our patients about DCIS, its heterogeneity, the benefits and risks of all the treatment options, and present the data in an unbiased way to help each patient make a decision that is right for her. We also must better understand the biology of the disease to make meaningful strides in how we manage the disease clinically. This requires access to well-annotated biospecimens for research. Moreover, encouraging the development and characterization of induct transplanted models, transgenic, and knockout mice models of DCIS may allow study of in situ cancer at all stages of progression to invasive cancer.

During the discussion sessions, it became clear that we must develop a common, standardized language to communicate about DCIS. Some debate exists as to whether or not “carcinoma” should be included in the naming of DCIS. If different clinicians speak about DCIS using different terms and with different perceptions of risks, then how do we expect our patients to have a clear understanding of their disease? We need to ensure that we are presenting the disease using a shared language. This language will be informed by the biological advances as they come, but in the meantime, it is important to remain consistent in our presentation and treatment discussions.

Finally, continuing to seek out and participate in regional, national, and international collaborations is another crucial step to furthering our understanding and our patients’ understanding of DCIS. There are a number of DCIS collaborative group clinical trials available, and we should encourage patients to consider these options. The trials are a result of institutional collaborations both nationally and internationally, and results from these trials have the potential to change the way we treat people with DCIS. Unraveling the biological drivers of cancer progression in DCIS and applying this knowledge to refine patient care exemplifies the goal of precision therapy. Combining biologic insights with improved strategies to elicit a patients’ treatment preferences will allow for the mitigation of both over-treatment and undertreatment. It is hoped that the lessons learned in DCIS research will inform an overall framework of how to rationally address the issues posed by cancer screening and early detection, not only for breast cancer but for other screen-detected cancers.

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