Introduction

Aromatase inhibitors (AIs) are critical to the management of women with hormone receptor-positive breast cancer (BC). Their use has evolved to include premenopausal women with high-risk BC in combination with ovarian suppression [1, 2]. Whether administered as monotherapy, sequential therapy, or extended therapy, AIs favorably impact disease free survival [3]. However, AIs are also associated with a number of toxicities, of which arthralgia (AIA) is among the most common and significant.

The broad range of reported incidence of AIA (5–47%) may be attributed to a lack of uniformity in diagnostic criteria to define the condition and casual reporting [4]. In two studies specifically designed to identify AIA, the incidence of AIA was consistently reported near 50%.

Genomic risk prediction of aromatase inhibitor-related arthralgia in patients with breast cancer using a novel machine-learning algorithm

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Keywords
Aromatase, arthralgia, breast cancer, estrogen, joint pain, SNP

Abstract

Many breast cancer (BC) patients treated with aromatase inhibitors (AIs) develop aromatase inhibitor-related arthralgia (AIA). Candidate gene studies to identify AIA risk are limited in scope. We evaluated the potential of a novel analytic algorithm (NAA) to predict AIA using germline single nucleotide polymorphisms (SNP) data obtained before treatment initiation. Systematic chart review of 700 AI-treated patients with stage I-III BC identified asymptomatic patients (n = 39) and those with clinically significant AIA resulting in AI termination or therapy switch (n = 123). Germline DNA was obtained and SNP genotyping performed using the Affymetrix UK BioBank Axiom Array to yield 695,277 SNPs. SNP clusters that most closely defined AIA risk were discovered using an NAA that sequentially combined statistical filtering and a machine-learning algorithm. NCBI PhenGenI and Ensemble databases defined gene attribution of the most discriminating SNPs. Phenotype, pathway, and ontologic analyses assessed functional and mechanistic validity. Demographics were similar in cases and controls. A cluster of 70 SNPs, correlating to 57 genes, was identified. This SNP group predicted AIA occurrence with a maximum accuracy of 75.93%. Strong associations with arthralgia, breast cancer, and estrogen phenotypes were seen in 19/57 genes (33%) and were functionally consistent. Using a NAA, we identified a 70 SNP cluster that predicted AIA risk with fair accuracy. Phenotype, functional, and pathway analysis of attributed genes was consistent with clinical phenotypes. This study is the first to link a specific SNP/gene cluster to AIA risk independent of candidate gene bias.
Therapy-Related Arthralgia Prediction

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Patients and Methods

After Institutional Review Board permission was obtained, a systematic chart review of women enrolled in The Columbus Breast Cancer Tissue Bank was completed to identify patients receiving first-line adjuvant treatment with third-generation AIs (anastrozole, exemestane, letrozole) for at least 1 month to treat stage I-III estrogen receptor-positive BC between 2003 and 2012. Clinical efficacy of endocrine therapy was not captured in this investigation. Concurrent gonadotropin-releasing (GnRH) agonist therapy, radiation therapy, prior tamoxifen use, and/or chemotherapy were allowed. Patients with metastatic disease or active autoimmune or inflammatory joint disease were excluded. Patients were divided into two groups: those with clinically significant AIA (defined as grade 2 or above by NCI-CTCv4 criteria) and/or requiring modification or termination of AI therapy, and those without any reported clinical signs or symptoms of AIA.

Germline DNA was extracted from mononuclear cells at the Human Cancer Genetics Sample Bank, The Ohio State University, according to previously published protocol [14]. DNA was quantitated using PicoGreen and 200 ng of each DNA aliquoted into 96-well plates. After sample elimination for poor quality DNA, SNP genotyping was performed along with appropriate controls. DNA amplification, fragmentation, and hybridization to Axiom UK Biobank genotyping arrays (Affymetrix, Santa Clara, CA) was completed using Axiom Reagent Kits; hybridization, ligation, washing, staining, and scanning of the arrays was completed on the GeneTitan MC instrument (Affymetrix). Initial plate QC was performed using Affymetrix Genotyping Console Software, and genotype calling done using Affymetrix Power Tools (v1.15.0) with the Axiom GT1 algorithm, which is a modified version of the BRLMM-P algorithm that adapts generic prior cluster positions to the data using an EM algorithm.

Analysis

To establish the parameters by which fold-change thresholds were maximized to identify distinguishing SNPs, we considered SNPs in 3 groups: those uniquely associated with the AIA-positive group, the AIA-negative group, and those predominantly, but not uniquely associated with either group. We then assigned arbitrary numerical identifiers of 1, 2, or 3 to each group for parameterization and found that the maximum fold-change defining a signal was \( \log_2 (3/1) = 1.59 \). Figure 1 shows the flowchart of the methodology, which is composed of the following steps:

Analysis of the discriminatory power of the SNPs

To identify the discriminatory power of those SNPs that were differentially noted between the AIA-positive and -negative cohorts, we utilized a machine-learning algorithm in which we combined fold change and Fisher’s ratio. We defined the Fisher’s ratio for a SNP \( j \) in a two-class classification problem, \( c_1, c_2 \) as:

\[
\text{FR}_{(c_1,c_2)} = \frac{(\mu_{c_1} - \mu_{c_2})^2}{\sigma_{c_1}^2 + \sigma_{c_2}^2}
\]

where \( \mu_{c_1}, \mu_{c_2} \) are measures of the center of the distribution (means) of prognostic variables \( j \) in classes 1 and 2 (AIA and no AIA), and \( \sigma_{c_1}^2, \sigma_{c_2}^2 \) are measures of the dispersion (variance) within these classes. This method is particularly effective for identifying prognostic/predictive variables that separate the classes further apart and are very homogeneous within classes (low intraclass variance).
We considered the 1% tail of the fold change (over and underexpressed SNPs), providing a final set of 436 high discriminatory SNPs with fold change in the interval $[-0.31, 0.22]$ and Fisher's ratio greater than 1.5.

Finding the small-scale SNPs signature

Once we identified those SNPs differentiating AIA-positive and -negative patients, we ranked them in decreasing order based on their discriminatory power. Hypothesizing that optimization of predictive risk determination is most accurately the consequence of a collective effect from a cluster of SNPs, we then sought to identify the smallest aggregate of SNPs with the highest prognostic accuracy using an algorithm based on recursive elimination of lower discriminatory SNPs. Our analysis was based on the fact that high discriminatory variables served to span the main features of the classification (AIA), while the variables with lowest discriminatory ratios were of such granularity as to not markedly contribute to being informative as to differential risk. This method determined the minimum amount of high-frequency details required to optimally discriminate between classes. The predictive accuracy estimation was based on Leave-One-Out-Cross-Validation (LOOCV) [10,11] given our goal to estimate how accurately the predictive model (classifier) would perform for future samples with an unknown AIA status.

Stability analysis of the small-scale signature

By random 75–25 hold-out experiments, we next evaluated the stability of the small-scale signatures’ predictive accuracy (i.e., AIA contributions) found via LOOCV when the number of training samples was decreased. Due to absence of a totally independent clinical data set, the minimum-scale signature was read in the training dataset for training (75% of the whole set) and applied for blind validation in the validation set (25%). The cumulative distribution function of the small-scale predictive accuracies found in different hold-outs was finally presented and accounted for the variability in its predictive accuracy with partial information. An additional statistical analysis was performed to provide the minimum, maximum, and median bounds that could be expected in an independent dataset. Figure 2 shows the cumulative probability function of the predictive accuracy of the small-scale signature obtained after 5000 random simulations.

Random sampling of high predictive SNPs equivalent networks

We next used a random sample to find other networks of highly discriminatory, prognostic SNPs. As the prior sampling probability of any individual SNP was considered to be proportional to its Fisher’s ratio, we preferentially sampled the most discriminatory SNPs. To look at the impact of SNP synergism in enhancing risk prediction, we developed the most discriminatory networks and analyzed the posterior sampling frequencies of the main prognostic variables involved in each network. The analysis was completed by establishing the correlation network among the most discriminatory SNPs. The network was built using the maximum spanning tree algorithm (an acyclic graph that maximizes the value of the edges) and the Pearson correlation coefficient to identify those SNPs that showed the maximum positive and negative Pearson coefficient.

Gene attribution and functional analysis

We employed two different databases, NCBI Phenotype-Genotype Integrator (NCBI PheGenI) [15] and Ensemble
release 88 [16] using Genome assembly: GRCh38.p9, to gather the genes and their functional consequences associated with the top 70 most discriminatory SNPs (not shown).

To assess the functional validity of the SNP-associated genes, we performed a comprehensive literature review of each gene associated with the most predictive SNPs, and summarized all relevant phenotypic attributions linking them to the following phenotypes: “arthralgia,” “synovial,” “arthritis,” “rheumatoid,” “joint,” “pain,” “sensitization,” and “nociception.” This was done via an ‘undirected’ review of all publications citing each gene, which was accessed via GeneAnalytics GeneCards’ [17] “Publications” section for each gene of interest [18].

We next determined the presence of any linkage disequilibrium (LD) among the 70 most discriminatory SNPs, as it is plausible that SNPs that individually have marginal influence on arthralgia risk-associated genes may be inherited in linkage, and together as a group, synergistically increase arthralgia risk. Linkage disequilibrium was assessed by querying the 70 SNPs using Broad Institute’s SNP Annotation and Proxy Search (SNAP) tool in the CEU (Utah residents with Northern and Western European ancestry) with $R^2$ threshold of 0.8 and distance limit of 500 [19, 20].

As a final analysis of functional and phenotypic relevance of the gene set found to be influenced by the 70 SNPs, we applied an undirected assessment using GeneAnalytics software. We reasoned this would provide a relevance validation, but acknowledge the risk of overinterpretation of a small input. This program provides a semiquantitative output of gene cluster relationships relative to disease, pathway, and ontologic relationships. Relationships with “medium” and “high” matched scores for the diseases and pathways were reviewed.

**Results**

Systematic chart review of 700 AI-treated patients with stage I-III BC identified asymptomatic patients ($n = 39$) and those with clinically significant AIA resulting in AI termination or therapy switch ($n = 123$). There were no significant demographic or disease differences between the AIA-positive and AIA-negative cohorts. Patients were similar in age (controls: 58.6 years, cases: 58.2 year), stage at diagnosis, and estrogen receptor status (Table 1). After sample elimination for poor-quality DNA, SNP genotyping was performed for 123 AIA positive and 39 AIA negative patients.

**Discriminatory SNPs**

Germline DNA was obtained and SNP genotyping performed to yield 695,277 SNPs. The analysis confirmed the importance of the main discriminatory SNPs (Table 2). Using the filtering sequence, after we determined and rank ordered the most discriminatory SNPs ($n = 400$), we identified the smallest number of SNPs that were most predictive of risk using the LOOCV algorithm described above. This method’s overall predictive accuracy was calculated by iterating all of the filtered samples. We found that a signature consisting of 70 specific SNPs had the highest predictive accuracy of 75.93% (Table 2). Analysis of the SNP signature’s predictive accuracy (Figure 2) demonstrated a median accuracy and true positive rate of 75.6% with an interquartile range of 7.3% and a true negative rate of 76.2%. The predictive accuracy’s mean and standard deviation was 74.6% and 5.8%, respectively. The minimum and maximum accuracy of these random holds was 54% and 100%, which implies that the minimum size signature of SNPs is quite stable. While we

| Table 1. Comparative tumor characteristics of patients in the control (no AIA) and cases (clinically significant AIA) arms. |
|-----------------------------------------------|
| Controls ($N = 39$) (%) | Cases ($N = 123$) % |
|----------------------|---------------------|
| Stage I              | 22 (56)             | 55 (45)            |
| Stage II             | 14 (36)             | 54 (44)            |
| Stage III            | 3 (8)               | 14 (11)            |
| ER or PR positive/HER2 negative | 33 (85) | 104 (85) |
| ER or PR positive/HER2 positive | 6 (15) | 19 (15) |
Table 2. The seventy SNPs that defined AIA risk within the study population, and their possible functional relevance.

| SNP          | SNP Consequence                      | Impacted Gene | Gene’s Relevant | Evidence/Comments                                                                 |
|--------------|--------------------------------------|---------------|-----------------|-----------------------------------------------------------------------------------|
| rs72765615   | Intron Variant, 3’ UTR Variant       | CHD2          | –               |                                                                                   |
| rs17149310   | Downstream Gene Variant              | CFAP77        | –               |                                                                                   |
| rs8028334    | Intron variant                       | IL16          | RA pathophysiology | Differentially elevated in synovial fluid from RA patients [25, 26, 44-48] and mediates chemoattraction of CD4+ cells to synovial tissue [25, 26, 49, 50]. However, IL16 correlation with clinical disease activity has been conflicting[45, 46]. |
| rs12004732   | Intron variant                       | PLAA          | RA pathophysiology | Detected in high levels in RA synovial fluid [22] and may have inflammatory roles [51]. Intrathecal injection in rabbit joints results in inflammatory arthritis[21] |
| rs2883917    | Intron Variant                       | NR3C2         | - Pain sensitization - Fibromyalgia | May be promote visceral hypersensitivity[36], and be implicated in pathophysiology of fibromyalgia[37] |
| rs61363926   | Noncoding Transcript Exon Variant    | BANF1P2       | –               |                                                                                   |
| rs56335940   | Intron Variant                       | LINC00882     | –               |                                                                                   |
| rs3749817    | Missense                             | FSTL4         | –               |                                                                                   |
| rs879605     | Intron variant, upstream variant 2KB | LTBR          | RA pathophysiology | Induces RA synovial fibroblast proliferation and expression of inflammatory elements [28,29]. Associated with pain and disability in RA patients [27] |
| rs879605     | Intron variant, upstream variant 2KB | SCNN1A        | –               |                                                                                   |
| rs986324     | Intron variant                       | PTCHD1-AS     | –               |                                                                                   |
| rs7017819    | Intron variant, Noncoding Transcript Exon Variant | RP1L1 | – |                                                                                   |
| rs12799692   | Intron variant                       | OPCML         | –               | OPCML may bind opioids [52]. It is also a tumor suppressor in and may be a marker of several types of tumors[53, 54] |
| rs4394668    | Intron variant                       | DHRS3         | –               |                                                                                   |
| rs10996945   | Intron variant                       | CTNNA3        | –               |                                                                                   |
| rs10996945   | Intron variant                       | LOC105378340 | –               |                                                                                   |
| rs705226     | Intron variant                       | LOC105374060 | –               |                                                                                   |
| rs73042968   | Intron Variant                       | FBLN2         | Breast cancer pathophysiology | Loss of FBLN2 expression is associated with breast cancer progression [55] |
| rs1546734    | Intron Variant                       | LOC105377150 | –               |                                                                                   |
| rs17270243   | Intron Variant                       | RORA-AS1      | - Breast cancer pathophysiology - Estrogen metabolism | - May suppress breast tumor invasion by inducing SEMA3F[56]. - [Conflicting] Is a transcriptional regulator of aromatase [57]. Activates aromatase expression in breast cancer cells, likely contributing to proliferation [58] |
| rs17270243   | Intron Variant                       | RORA*         | - Breast cancer pathophysiology - Estrogen metabolism |                                                                                   |
| rs5760686    | Intron Variant                       | SGSM1         | –               |                                                                                   |
| rs9907168    | Intron Variant                       | CDC42EP4      | –               |                                                                                   |
| rs76098632   | Intron Variant                       | FBXL17        | Breast cancer marker | May be a potential biomarker for breast cancer therapy[59] |
| rs2243511    | Intron Variant                       | TMEM508       | –               |                                                                                   |
### Table 2 (Continued)

| SNP          | SNP Consequence | Impacted Gene | Gene’s Relevant Phenotype | Evidence/Comments                                                                 |
|--------------|-----------------|---------------|---------------------------|----------------------------------------------------------------------------------|
| rs2243511    | Intron Variant  | IFNGR2        | RA pathophysiology        | - Significant differences in blood mononuclear cell expression of IFNGR2 was seen in African American RA patients with erosion and those with no erosion [30] |
| rs1012629    | Intron Variant  | PTPRK         | - RA pathophysiology      | - Knocing down its encoded receptor impairs migration and invasiveness of RA fibroblast-like synoviocytes (FLS), which otherwise progress to destroy cartilage and bone. This receptor mediates inflammatory signaling of TGF-Beta in RA FLS [60]. |
|              |                 |               | - Breast cancer pathophysiology |                                                                                 |
| rs322960     | Intron Variant  | TRPV3         | Pain sensitization        | - TRPV3 activation senses peripheral pain [24]. It accumulates in peripheral nerves and dorsal root ganglion after injury [23]. |
| rs3743160    | Intron Variant, 5′ UTR Variant | SLC28A1 | Breast cancer pathophysiology | - SLC28A1 expression may be implicated breast cancer cell responsiveness to chemotherapy [62, 63] |
| rs797818     | Intron Variant  | SEMA3A        | - RA pathophysiology      | - Reduced SEMA3A expression in human synovial tissues was associated with RA disease activity [31] |
|              |                 |               | - OA pathophysiology      | - SEMA3A expression is elevated in osteoarthritic cartilage, and inhibits VEGF’s effects [64] |
| rs11670284   | Intron Variant  | NLRP13        | –                         | *Questionable relevance:                                                                |
| rs2808787    | Intron Variant  | COL27A1       | –                         | - The temporal association and location of COL27A1 encoded collagen during calcification/transition of cartilage to bone suggests that the collagen is involved in the process. However, no specific roles have been elucidated [65]. |
| rs2215016    | Intron Variant  | RGS6          | –                         | - A SNP in the region of COL27A1 (rs946053) occurred significantly more in a sample of patients with Achilles Tendinopathy, than in the control group [66]. |
| rs3766160 and rs3820071 | Missense Variant | CELA2B | Poorly studied. | |
| rs10511813   | Upstream variant 2KB | LOC105376002 | –                         | |
| rs12127403   | Upstream Gene Variant | VHL | Poorly studied. | |
| rs10908495   | Missense, Noncoding Transcript Variant | GLMP | –                         | |
| rs11683506   | Intron Variant  | SMARCAL1      | –                         | *Questionable relevance:                                                                |
| rs6081792    | Intron Variant, Upstream Gene Variant | RIN2 | –                         | - Deficiency of encoded protein causes a congenital syndrome that includes severe joint hyperlaxity and scoliosis [67]. |
| rs1047312    | 3′ UTR Variant  | SULT1C2       | –                         | |
| rs17011869   | Intron Variant  | CNTNAP5       | –                         | |
| rs8183999    | Intron Variant  | LINC00922     | –                         | |
| SNP          | SNP Consequence                      | Impacted Gene        | Gene’s Relevant Phenotype Attributions | Evidence/Comments                                                                 |
|-------------|--------------------------------------|----------------------|---------------------------------------|-----------------------------------------------------------------------------------|
| rs11586047  | Intron Variant                       | LOC105371436         | –                                     |                                                                                  |
| rs28964     | Intron Variant                       | SPACA3               | –                                     |                                                                                  |
| rs10900269 and rs11239786 | Intron Variant, Noncoding Transcript Variant | BMS1               | –                                     |                                                                                  |
| rs11600377  | Noncoding Transcript, Exon Variant   | MRGPRF-AS1           | –                                     |                                                                                  |
| rs11600377  | Upstream Gene Variant                | MRGPRF              | –                                     |                                                                                  |
| rs62525208  | Intron Variant                       | C8orf37-AS1          | –                                     |                                                                                  |
| rs61782448  | Intron Variant                       | PLEKHM2              | –                                     |                                                                                  |
| rs77413365  | Intron Variant                       | GRIA1               | Inflammatory pain sensitization       |                                                                                  |
| rs1280408   | Intron Variant                       | CGNL1               | –                                     |                                                                                  |
| rs13013882  | Splice Region Variant, Synonymous Codon | MROH2A           | –                                     |                                                                                  |
| rs17599018  | Intron Variant                       | GPM6A               | Pain sensitization                    |                                                                                  |
| rs2269767   | Intron Variant                       | UBFD1               | Inflammatory antagonist               |                                                                                  |
| rs7024415   | Downstream Gene Variant              | ENSG00000253400     | –                                     |                                                                                  |
| rs1546734   | Intron Variant                       | ENSG00000242120     | –                                     |                                                                                  |
| rs4785496   | Upstream Gene Variant                | ENSG00000260605     | –                                     |                                                                                  |

Seventy SNPs were identified which were associated with AIA risk. In addition to listing those SNPs, the consequence of each SNP is defined as is the gene most impacted by the mutation. Using literature mining as described in the Methods section, we determined the potential implication of genes relative to the development of AIA.
**Figure 3.** Color visualization delineating the relationship of the 70 predictive SNPs and their functional relevance. (A) Correlation tree of the most discriminatory SNPs. This tree is built using the minimum spanning tree algorithm using the Pearson correlation coefficient. The algorithm looks for the maximum absolute values of the Pearson correlation coefficient (positive and negative correlations) within the set of most discriminatory SNPs. This hierarchical figure describes the strength of relationships between SNPs and how each SNP relates to the others in the cluster. (B) Associated phenotypes include similar phenotypes (RA, pain, inflammation and those associated with the tumor diagnosis).
cannot analyze how this predictive signature would vary when new patient data is acquired, we hypothesize the result will be similar to that described by the hold-out experiments.

The four most discriminating SNPs according to their sampling frequency as established by the random sampler were rs1462506, rs17149310, rs28839197, and rs10778060. The correlation network in Figure 3 shows that the header SNP in the graph (rs7276615) is weakly, negatively correlated with rs11586047, rs6195687, rs10916270, and rs1462506. The main tree is developed under rs11586047, related with rs11586047, rs6195687, rs10916270, and SNPs in the graph (rs7276615) are weakly, negatively correlated.

Functional validity and gene attribution

To assess the potential functional validity of the 70 most predictive SNPs, we assessed gene attribution for each SNP and found that 57 genes were associated with the 70 SNPs of interest, with four genes linked to consequences from two or more SNPs (rs12004732 and rs7863476, variants of PLAA; rs322960 and rs60292929, variants of TRPV3; rs3766160 and rs3820071, variants of CELA2B, and rs10900269 and rs11239786, variants of BMS1). Assuming that an increase in the ratio of SNP to gene could suggest a more influential impact of arthralgia risk prediction by that gene, we evaluated the relationship between the aforementioned genes and arthralgia-related phenotypes.

Following the search strategy defined above, we noted that both PLAA and TRPV3 had relevant associations with arthralgia phenotypes. PLAA is implicated in inflammatory pathways of synovial cells from Rheumatoid Arthritis (RA) patients [21, 22], and caused inflammatory arthritis when injected into rabbit knee joints [21]. TRPV3 accumulates in peripheral nerves and dorsal root ganglion [23], and its activation is implicated in sensing peripheral pain [24]. Associations of CELA2B are poorly understood. Lastly, there is no primary data to suggest a direct relationship between BMS1, arthralgias or BC, although BMS1 is also poorly studied. The potential synergistic impact of these genes on AIA risk requires additional study.

In addition to the four genes above, 19 of the other 57 genes associated with the top 70 SNPs were linked to phenotypes involving BC, estrogen metabolism, and significantly, arthralgia. Some of these arthralgia associations include IL16 (was significantly elevated in synovial fluid of RA patients and mediated chemotraction of CD4 + cells into synovial tissues [25,26]), LTBR (its expression in RA patients’ synovium positively associated with pain and disability [27], and may be implicated in RA synovial fibroblast proliferation and expression of inflammatory elements [28, 29]), IFNGR2 (joint erosion, joint space narrowing, and disease progression was significantly associated with differential expression of IFNGR2 in blood mononuclear cells of African American RA patients with radiographic erosion compared to patients with no erosion [30]), SEMA3A (expressed in synovial tissues and associated with RA [31]), GRIA1 (implicated in inflammatory pain and central sensitization of dorsal horns [32–34]), GPM6A (associated with endocytosis of Mu-type opioid receptors in neuronal cortical cell cultures [35]), NR3C2 (may promote visceral hypersensitivity [36] and may be implicated in pathophysiology of fibromyalgia [37]), and lastly, UBD1 (implicated as a regulator of NFkB pathway [38]) (Table 2).

Linkage disequilibrium

The LD analysis found 3 sets of SNPs among the top 70 SNP to be in LD. The first LD pair included rs3766160 and rs3820071, missense mutations of CELA2B, a poorly characterized gene. The second pair includes LD between rs12127403 (Upstream Gene Variant of VHLL, which competitively prevents degradation of HIF-alpha [39]) and rs10908495 (a variant of GLMP, another poorly characterized gene). The last pair consisted of rs3011665 and rs981360, which are not known to consequence any specific genes.

Discussion

AIA is a prevalent and disrupting toxicity of AIs, impacting quality of life, adherence and clinical outcomes [40]. As with other regimen-related toxicities, the risk for AIA is not consistent among patients receiving identical treatments for the same disease. The ability to prospectively differentiate patients at risk for AIA would be desirable at a number of levels, but most importantly in providing actionable data that could inform clinician and patient decision-making, as well as leading to the design of intervention trials focused on high-risk individuals. Additionally, the identification of a predictive biomarker strongly associated with a clinically meaningful manifestation of AIA could provide a surrogate for its more accurate reporting.

To identify the SNPs of interest, we used a machine-learning algorithm for which the SNPs of interest were not pre-determined. Rather, employing a previously validated technique, we used a filtering step to create a hierarchical list of SNPs most associated with the AIA phenotype while simultaneously eliminating SNPs that were simply genomic “noise” [13]. We were able to reduce nearly 400,000 SNPs by three logs to approximately 450 SNPs. We then asked which SNPs were most predictive as a
‘team’ and, using a method in which we sequentially tested every combination of SNPs, identified 70 SNPs that collectively predicted AIA with fair accuracy (75.93%).

Attempts to determine AIA risk-based strictly on demographic features have been only marginally successful. We found no differences in the study cohorts’ clinical or demographic characteristics. The application of genomic markers, particularly SNPs, as a means to assess toxicity risk associated with cancer treatment regimens has been studied broadly, but with wide ranging and often inconsistent results. In general, two approaches have been used: candidate gene or SNP studies and GWAS. Investigations of genomic risk factors for AIA have exclusively depended on candidate gene/SNP identification.

Since CYP19A1 codes for the aromatase enzyme in postmenopausal women, it has been an obvious candidate gene for AIA risk prediction. While three studies have studied polymorphisms associated with CYP19A1 relative to AIA and found an association, there is variability in the SNPs reported. Other polymorphisms associated with estrogen and vitamin D metabolism have also been targeted. While analyzing such SNPs for risk prediction, García-Giralt and her colleagues introduced multiplicative terms into their analysis, thus providing a conceptual basis for synergism in contributing to AIA risk. In a related investigation, Lintemann et al. reported that a SNP associated with the osteoprotegrin gene was associated with adverse symptoms (hot flashes and pain) in patients treated with AIs.

A drawback of candidate gene studies is that they limit discovery of phenotype-associated genes or SNPs that may not be obvious to investigators deciding on targets. Analogous to trying to describe a landscape in the dark by shining a flashlight with a narrow beam, candidate gene studies may miss important features. Consequently, we took an analytical approach that differed from conventional paradigms in three important ways: (1) we did not mandate a threshold gene expression (SNP) level change for inclusion; (2) we evaluated simultaneous expression of SNP profiles (clusters); (3) the predictive clusters that evolved were driven by their collective and hierarchical relationships with the study groups rather than dependent on preconceptions of an expected result. Importantly, we were able to confirm the relevance of the discovered clusters by confirming their fit into known ontological pathways.

To determine the functional validity of the SNPs in the cluster, we attributed SNPs to their related genes (n = 57) and evaluated relevance to several phenotypes that we thought may be expressed in the study cohort. Many individual SNPs were functionally specific for arthralgia-like disease phenotypes. Others were associated with estrogen metabolism – a finding that supports the hypotheses suggested by the CYP21A candidate studies. None of the SNPs in this cluster were attributable to CYP21A genes, which may be a component of faults in SNP and gene designation. Genes associated with pain sensitization were notable, a finding theoretically similar to the interests of Lintemann et al. Interestingly, genes associated with RA pathophysiology were relatively high in the cluster, suggesting a common biological pathway with AIA and in congruence with other findings implicating an inflammatory mechanism. One SNP in the cluster was closely aligned with inflammation.

Although we included internal cross-validation in this study, the investigation was limited by small sample size and our inability to have an independent validation cohort. We believe that increasing the training sample will result in a more robust predictive accuracy. An independent validation cohort will be critical to understanding the true clinical meaningfulness and translatability of our findings. A prospective study is currently underway to address these shortcomings.

Nonetheless, we believe that this trial demonstrates the potential utility of an undirected, machine-learning approach in the development of a predictive test for AIA risk. Such a personalized model in which at-risk patients are identified prior to therapy start may help to minimize toxicity by prompting the early institution of preventative, therapeutic, or alternative interventions, and thus improve treatment adherence and disease outcomes.

**Conflict of Interest**

Disclosures outside the submitted work are noted. Stephen Sonis, Ohio State University (grants), Primary Endpoint Solutions LLC (equity), Biomodels LLC (employee), Inform Genomics (equity); Juan Luis Fernández-Martínez Primary Endpoint Solutions LLC (consultant); Ana Cernea Primary Endpoint Solutions LLC (consultant); Enrique J. de Andrés-Galiana Primary Endpoint Solutions LLC (consultant).

**References**

1. Pagani, O., M. M. Regan, and P. A. Francis. 2014. Investigators TaS, group IBCS. Exemestane with ovarian suppression in premenopausal breast cancer. N. Engl. J. Med. 371:1358–1359.

2. Francis, P. A., M. M. Regan, and G. F. Fleming. 2015. Adjuvant ovarian suppression in premenopausal breast cancer. N. Engl. J. Med. 372:1673.

3. Aydiner, A. 2013. Meta-analysis of breast cancer outcome and toxicity in adjuvant trials of aromatase inhibitors in postmenopausal women. Breast (Edinburgh, Scotland) 22:121–129.
4. Niravath, P. 2013. Aromatase inhibitor-induced arthralgia: a review. Ann. Oncol. 24:1443–1449.
5. Henry, N. L., J. T. Giles, D. Ang, L. Henry, Jon. T. Giles, D. Ang, M. Mohan, et al. 2008. Prospective characterization of musculoskeletal symptoms in early stage breast cancer patients treated with aromatase inhibitors. Breast Cancer Res. Treat. 111:365–372.
6. Crew, Katherine. D., H. Greenlee, J. Capodice, G. Raptis, L. Braffman, D. Fuentes, A. Sierra, Dawn. L. Hershman, et al. 2007. Prevalence of joint symptoms in postmenopausal women taking aromatase inhibitors for early-stage breast cancer. J. Clin. Oncol. 25:3877–3883.
7. Partridge, A. H., A. LaFountain, E. Mayer, B. S. Taylor, E. Winer, and A. Asnis-Alibuzek. 2008. Adherence to initial adjuvant anastrozole therapy among women with early-stage breast cancer. J. Clin. Oncol. 26:556–562.
8. Mao, Jun. J., C. Stricker, D. Bruner, S. Xie, Marjorie. A. Bowman, John. T. Farrar, Brandon. T. Greene, A. DeMichele, et al. 2009. Patterns and risk factors associated with aromatase inhibitor-related arthralgia among breast cancer survivors. Cancer 115:3631–3639.
9. Cecil, R. L., and B. H. Archer. 1925. Arthritis of the menopause: a study of fifty cases. J. Am. Med. Assoc. 84:75–79.
10. Lintermans, A., K. Van Asten, L. Jongen, T. Van Brussel, A. Laenen, J. Verhaeghe, et al. 2016. Genetic variant in the osteoprotegerin gene is associated with aromatase inhibitor-related musculoskeletal toxicity in breast cancer patients. Eur. J. Cancer (Oxford, England: 1990)56:31–36.
11. Wang, J., L. Kangping, Y. Song, S. Zhao, W. Ma, Q. Xuan, D. Tang, H. Zhao, L. Liu, Q. Zhang, et al. 2015. RANKL and OPG Polymorphisms Are Associated with Aromatase Inhibitor-Related Musculoskeletal Adverse Events in Chinese Han Breast Cancer Patients. PLoS ONE 10:e0133964.
12. Garcia-Giralt, N., M. Rodriguez-Sanz, D. Prieto-Alhambra, S. Servitja, E. T.-d. Pliego, S. Balcoks, J. Albanell, D. Grinberg, A. Diez-Perez, I. Tusquets, X. Nogués, et al. 2013. Genetic determinants of aromatase inhibitor-related arthralgia: the B-ABLE cohort study. Breast Cancer Res. Treat. 140:385–395.
13. Saligan, L. N., I. L. Fernández-Martínez, E. J. deAndrés-Galiana, and S. Sonis. 2014. Supervised classification by filter methods and recursive feature elimination predicts risk of radiotherapy-related fatigue in patients with prostate cancer. Cancer Inform. 13:141–152.
14. Miller, S. A., D. D. Dykes, and H. F. Polesky. 1988. A simple salting out procedure for extracting DNA from human nucleated cells. Nucleic Acids Res. 16:1215.
15. Ramos, Erin. M., D. Hoffman, Heather. A. Junkins, D. Maglott, L. Phan, Stephen. T. Sherry, M. Feolo, Lucia. A. Hindorff, et al. 2014. Phenotype-Genotype Integrator (PheGenI): synthesizing genome-wide association study (GWAS) data with existing genomic resources. Eur. J. Hum. Genet. 22:144–147.
16. Yates, A., M. Wasiu Akanni, R. Amode, D. Barrell, K. Bills, D. Carvalho-Silva, C. Cummins, P. Clapham, et al. 2016. Ensembl 2016. Nucleic Acids Res. 44(D1):D710–716.
17. Fuchs, S. B.-A., I. Lieder, G. Stelzer, Y. Mazor, E. Buzhor, S. Kaplan, Y. Bogoch, I. Plaschkes, et al. 2016. GeneAnalytics: an integrative gene set analysis tool for next generation sequencing, rnaseq and microarray data. OMICS 20:139–151.
18. Stelzer, G., N. Rosen, I. Plaschkes, S. Zimmerman, M. Twik, S. Fishilevich, T. I. Stein, R. Nudel, I. Lieder, Y. Mazor, S. Kaplan, D. Dahary, D. Warshawsky, Y. Guan-Golan, A. Kohn, N. Rappaport, M. Safran, D. Lambert, et al. 2016. The GeneCards Suite: From Gene Data Mining to Disease Genome Sequence Analyses. Curr. Protoc. Bioinformatics. 54:1.30.31-31.30.33.
19. Johnson, A. D., R. E. Handsaker, S. L. Pulit, M. M. Nizzari, C. J. O’Donnell, and P. I. de Bakker. 2008. SNAP: a web-based tool for identification and annotation of proxy SNPs using HapMap. Bioinformatics 24:2938–2939.
20. Frazer, Kelly. A., Dennis. G. Ballinger, David. R. Cox, David. A. Hinds, Laura. L. Stuve, Richard. A. Gibbs, John. W. Belmont, A. Boudreau, P. Hardenbol, Suzanne. M. Leal, S. Pasternak, David. A. Wheeler, Thomas. D. Willis, Y. Fuli, H. Yang, C. Zeng, Y. Gao, et al. 2007. A second generation human haplotype map of over 3.1 million SNPs. Nature 449:851–861.
21. Bomalaski, J. S., M. Fallon, R. A. Turner, S. T. Crooke, P. C. Meunier, and M. A. Clark. 1990. Identification and isolation of a phospholipase A2 activating protein in human rheumatoid arthritis synovial fluid: induction of eicosanoid synthesis and an inflammatory response in joints injected in vivo. J. Lab. Clin. Med. 116:814–825.
22. Goddard, D. H., S. L. Grossman, R. Newton, M. A. Clark, and J. S. Bomalaski. 1992. Regulation of synovial cell growth: basic fibroblast growth factor synergizes with interleukin 1 beta stimulating phospholipase A2 enzyme activity, phospholipase A2 activating protein production and release of prostaglandin E2 by rheumatoid arthritis synovial cells in culture. Cytokine 4:377–384.
23. Facer, P., Maria. A. Casula, Graham. D. Smith, Christopher. D. Benham, lain. P. Chessel, C. Bountra, M. Sinisi, R. Birch, P. Anand, et al. 2007. Differential expression of the capsacin receptor TRPV1 and related novel receptors TRPV3, TRPV4 and TRPM8 in normal human tissues and changes in traumatic and diabetic neuropathy. BMC Neurol. 7:11.
24. Bang, S., S. Yoo, T. J. Yang, H. Cho, and S. W. Hwang. 2010. Farnesyl pyrophosphate is a novel...
pain-producing molecule via specific activation of TRPV3. J. Biol. Chem. 285:19362–19371.

25. Franz, Juliane. K., Stefan. A. Kolb, Klaus. M. Hummel, F. Lahrtz, M. Neidhart, Wilhelm. K. Aicher, T. Pap, Renate. E. Gay, A. Fontana, S. Gay, et al. 1998. Interleukin-16, produced by synovial fibroblasts, mediates chemoattraction for CD4 + T lymphocytes in rheumatoid arthritis. Eur. J. Immunol. 28:2661–2671.

26. Warstat, K., M. Hoberg, M. Rudert, S. Tsui, T. Pap, B. Franz, Juliane. K., Stefan. A. Kolb, Klaus. M. Hummel, O'Rourke, Killian. P., G. O'Donoghue, C. Adams, H. Park, J.-S., M. Yaster, X. Guan, X. Ji-Tian, M.-H. Takagawa, S., F. Nakamura, K. Kumagai, Y. Nagashima, O'Rourke, Killian. P., G. O'Donoghue, C. Adams, H. Park, J.-S., M. Yaster, X. Guan, X. Ji-Tian, M.-H. Takagawa, S., F. Nakamura, K. Kumagai, Y. Nagashima, et al. 2008. LIGHT (TNFSF14), a novel mediator of bone resorption, is elevated in rheumatoid arthritis. Arthritis Rheum. 54:1451–1462.

30. Tang, Q., M. I. Danila, X. Cui, L. Parks, B. J. Baker, R. J. Reynolds, C. Raman, and K. C. Wansek, et al. 2015. Expression of interferon-gamma receptor genes in peripheral blood mononuclear cells is associated with rheumatoid arthritis and its radiographic severity in African Americans. Arthritis Rheumatol. (Hoboken, NJ). 67:1165–1170.

31. Takagawa, S., F. Nakamura, K. Kumagai, Y. Nagashima, Y. Goshima, and T. Saito. 2013. Decreased semaphorin3A expression correlates with disease activity and histological features of rheumatoid arthritis. BMC Musculoskelet. Disord. 14:40.

32. Park, J.-S., M. Yaster, X. Guan, X. Ji-Tian, M.-H. Shih, Y. Guan, Srinivasa. N. Raja, Y.-X. Tao, et al. 2008. Role of spinal cord alpha-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptors in complete Freund’s adjuvant-induced inflammatory pain. Mol. Pain. 4:67.

33. Vikman, K. S., R. H. Hill, E. Backstrom, B. Robertson, and K. Kristensson. 2003. Interferon-gamma induces characteristics of central sensitization in spinal dorsal horn neurons in vitro. Pain 106:241–251.

34. Choi, J. I., C. I. Svensson, F. J. Koehrn, A. Bhuskute, and L. S. Sorkin. 2010. Peripheral inflammation induces tumor necrosis factor dependent AMPA receptor trafficking and Akt phosphorylation in spinal cord in addition to pain behavior. Pain 149:243–253.
fibroblasts from rheumatoid arthritis patients differ in their regulation of IL-16 gene activity in comparison to osteoarthritis fibroblasts. Cell. Physiol. Biochem. 14(4–6):293–300.

45. Blaschke, S., H. Schulz, G. Schwarz, V. Blaschke, G. A. Muller, and M. Reuss-Borst. 2001. Interleukin 16 expression in relation to disease activity in rheumatoid arthritis. J. Rheumatol. 28:12–21.

46. Kageyama, Y., T. Ozeki, M. Suzuki, et al. 2000. Interleukin-16 in synovial fluids from cases of various types of arthritis. Joint, Bone, Spine 67:188–193.

47. Lard, L. R., B. O. Roep, R. E. Toes, and T. W. Huizinga. 2004. Enhanced concentrations of interleukin 16 are associated with joint destruction in patients with rheumatoid arthritis. J. Rheumatol. 31:35–39.

48. Luo, S.-X., S. Li, X.-H. Zhang, J.-J. Zhang, G.-H. Long, G.-F. Dong, S. Wei, Y. Deng, Y. Liu, J.-M. Zhao, X. Qin, et al. 2015. Genetic polymorphisms of interleukin-16 and risk of knee osteoarthritis. PLoS ONE 10:e0123442.

49. McFadden, C., R. Morgan, S. Rahangdale, D. Green, H. Yamasaki, D. Center, and W. Cruikshank, et al. 2007. Preferential migration of T regulatory cells induced by IL-16. J. Immunol. (Baltimore, Md: 1950) 179:6439–6445.

50. Wilson, K. C., D. M. Center, and W. W. Cruikshank. 2004. The effect of interleukin-16 and its precursor on T lymphocyte activation and growth. Growth Factors (Chur, Switzerland) 22:97–104.

51. Ribardo, D. A., J. W. Peterson, and A. K. Chopra. 2002. Phospholipase A2-activating protein—an important regulatory molecule in modulating cyclooxygenase-2 and tumor necrosis factor production during inflammation. Indian J. Exp. Biol. 40:129–138.

52. Duarte-Pereira, S., F. Paiva, V. L. Costa, J. Ramalho-Carvalho, J. Savva-Bordalo, Á. Rodrigues, F. R. Ribeiro, Vitor. M. Silva, J. Oliveira, R. Henriques, C. Jerónimo, et al. 2011. Prognostic value of opioid binding protein/cell adhesion molecule-like promoter methylation in bladder carcinoma. Eur. J. Cancer 47:1106–1114.

53. Cui, Y., Y. Ying, A. van Hasselt, K. M. Ng, Y. Jun, Q. Zhang, J. Jin, D. Liu, Johng. S. Rhim, S. Y. Rha, M. Loyo, Anthony, T. C. Chan, G. Srivastava, George. S. W. Tsao, Grant, C. Sellier, Joseph. I. Y. Sung, D. Sidransky, Q. Tao, et al. 2008. OPCLM is a broad tumor suppressor for multiple carcinomas and lymphomas with frequently epigenetic inactivation. PLoS ONE 3:e2990.

54. Zhou, F., M. Ma, G. Tao, X. Chen, W. Xie, Y. Wang, and X. Cao. 2014. Detection of circulating methylated opioid binding protein/cell adhesion molecule-like gene as a biomarker for ovarian carcinoma. Clin. Lab. 60:759–765.

55. Yi, C. H., D. J. Smith, W. W. West, and M. A. Hollingsworth. 2007. Loss of fibulin-2 expression is associated with breast cancer progression. Am. J. Pathol. 170:1535–1545.

56. Xiong, G., C. Wang, B. M. Evers, B. P. Zhou, and R. Xu. 2012. RORalpha suppresses breast tumor invasion by inducing SEMA3F expression. Can. Res. 72:1728–1739.

57. Sarachana, T., M. Xu, R. C. Wu, and V. W. Hu. 2011. Sex hormones in autism: androgens and estrogens differentially and reciprocally regulate RORA, a novel candidate gene for autism. PLoS ONE 6:e17116.

58. Odawara, H., T. Iwasaki, J. Horiguchi, N. Rokutanda, K. Hirooka, W. Miyazaki, Y. Koibuchi, N. Shimokawa, Y. Iino, I. Takeyoshi, N. Koibuchi, et al. 2009. Activation of aromatase expression by retinoic acid receptor-related orphan receptor (ROR) alpha in breast cancer cells: identification of a novel ROR response element. The Journal of biological chemistry. 284:17711–17719.

59. Xiao, G. G., B. S. Zhou, G. Somlo, et al. 2008. Identification of F-box/LLR-repeated protein 17 as a potential useful biomarker for breast cancer therapy. Cancer Genomics Proteomics 5(3–4):151–160.

60. Stanford, Stephanie. M., German. R. Aleman, B. B. Muench, C. Sacchetti, William. B. Kiosses, J. Sharma, Michael. F. Maestre, M. Bottini, T. Mustelin, David. L. Boyle, Gary. S. Firestein, N. Bottini, et al. 2016. TGFBeta responsive tyrosine phosphatase promotes rheumatoid synovial fibroblast invasiveness. Ann. Rheum. Dis. 75:295–302.

61. Sun, P. H., L. Ye, M. D. Mason, and W. G. Jiang. 2013. Protein tyrosine phosphatase kappa (PTPRK) is a negative regulator of adhesion and invasion of breast cancer cells, and associates with poor prognosis of breast cancer. J. Cancer Res. Clin. Oncol. 139:1129–1139.

62. Lane, J., T. A. Martin, C. McGuigan, M. D. Mason, and W. G. Jiang. 2010. The differential expression of hCNT1 and hENT1 i n breast cancer and the possible impact on breast cancer therapy. J. Exp. Ther. Oncol. 8:203–210.

63. Cano-Soldado, P., M. Molina-Arcas, B. Algueró, I. Larráyoz, M. Pilar Lostao, A. Grandas, F. Javier Casado. 2008. Compensatory effects of the human nucleoside transporters on the response to nucleoside-derived drugs in breast cancer MCF7 cells. Biochem. Pharmacol. 75:639–648.

64. Okubo, M., T. Kimura, Y. Fujita, S. Mochizuki, Y. Niki, H. Enomoto, Y. Suda, Y. Toyama, Y. Okada, et al. 2011. Semaphorin 3A is expressed in human osteoarthritic cartilage and antagonizes vascular endothelial growth factor 165-promoted chondrocyte migration: an implication for chondrocyte cloning. Arthritis Rheum. 63:3000–3009.

65. Hjorten, R., U. Hansen, Robert. A. Underwood, Helena. E. Telfer, Russell. J. Fernandes, D. Krakow, E. Sebald, S. 252

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Wachsmann-Hogiu, P. Bruckner, R. Jacquet, William J. Landis, Peter. H. Byers, and James. M. Pace. 2007. Type XXVII collagen at the transition of cartilage to bone during skeletogenesis. Bone 41:535–542.

66. Saunders, Colleen. J., L. van der Merwe, M. Posthumus, J. Cook, Christopher. J. Handley, M. Collins, Alison. V. September, et al. 2013. Investigation of variants within the COL27A1 and TNC genes and Achilles tendinopathy in two populations. J. Orthop. Res. 31:632–637.

67. Syx, D., F. Malfait, L. Van Laer, J. Hellemans, T. Hermanns-Lê, A. Willaert, A. Benmansour, A. De Paepe, and A. Verloes. 2010. The RIN2 syndrome: a new autosomal recessive connective tissue disorder caused by deficiency of Ras and Rab interactor 2 (RIN2). Hum. Genet. 128:79–88.

68. Dong, X., S. Han, M. J. Zylka, M. I. Simon, and D. J. Anderson. 2001. A diverse family of GPCRs expressed in specific subsets of nociceptive sensory neurons. Cell 106:619–632.

69. Shozu, M., S. Sebastian, K. Takayama, W.-T. Hsu, Roger. A. Schultz, K. Neely, M. Bryant, and Serdar. E. Bulun. 2003. Estrogen excess associated with novel gain-of-function mutations affecting the aromatase gene. N. Engl. J. Med. 348:1855–1865.

70. Demura, M., R. M. Martin, M. Shozu, S. Sebastian, K. Takayama, W.-T. Hsu, R. A. Schultz, K. Neely, M. Bryant, B. B. Mendonca, K. Hanaki, S. Kanzaki, D. B. Rhoads, M. Misra, S. E. Bulun. 2007. Regional rearrangements in chromosome 15q21 cause formation of cryptic promoters for the CYP19 (aromatase) gene. Hum. Mol. Genet. 16:2529–2541.