Evaluating the Role of Circulating MicroRNAs to Aid Therapeutic Decision Making for Neoadjuvant Chemotherapy in Breast Cancer

A Prospective, Multicenter Clinical Trial

Matthew G. Davey, MCh, MRCSI,* Maire Caitlin Casey, MRCSI, PhD,* Andrew McGuire, PhD, MRCS,* Ronan M. Waldron, MB, BCh,* Maxwell Paganga, BSc† Emma Holian, PhD,† John Newell, MS, PhD,† Helen M. Heneghan, PhD, FRCSL,* Ailbhe M. McDermott, PhD,* Maccon M. Keane, MD,‡ Aoife J. Lowery, PhD, FRCSL, Nicola Miller, PhD,* and Michael J. Kerin, MCh, FRCS§

Objective: To evaluate whether circulating micro ribonucleic acids (miRNAs) predict response to neoadjuvant chemotherapy (NAC) and inform decision-making in breast cancer patients.

Introduction: Deciphering response to NAC remains a challenge. Those unlikely to respond may benefit from NAC de-escalation before completion, while “responders” should complete treatment. Establishing biomarkers which identify response to NAC is imperative to personalize treatment strategies. miRNAs are small noncoding RNA molecules which modulate genetic expression. miRNAs are believed to inform response to NAC.

Methods: This prospective, multicenter trial (NCT01722851) recruited 120 patients treated with NAC across 8 Irish treatment sites. Pre-determined miRNAs were quantified from patient whole bloods using relative quantification polymerase chain reaction. Venous sampling was performed at diagnosis and midway during NAC. Trends in miRNA expression between timepoints were correlated with treatment response. Data analysis was performed using R 3.2.3.

Results: A total of 120 patients were included (median age: 55 years). Overall, 49.2% had luminal breast cancers (59/120), 17.5% luminal B (L/B), 20.8% triple negative disease (25/120), and 10.8% HER2+ (15/120). For patients with luminal breast cancers, reduced Let-7a predicted achieving a pathological complete response (pCR) (21/120) (P = 0.012), while reduced miR-145 predicted response to NAC in HER2+ (P = 0.033). For patients with luminal breast cancers, reduced Let-7a predicted achieving a pCR (P = 0.037) and reduced miR-145 predicted achieving a pCR to NAC in HER2+ (P = 0.027).

Conclusions: This study illustrates the potential value of circulating miRNA measurement in predicting response to NAC. Further interrogation of these findings may see miRNAs personalize therapeutic decision-making for patients undergoing NAC for early breast cancer.

Keywords: breast cancer, miRNA, neoadjuvant therapies, personalized medicine, precision oncology

Although breast cancer incidence is increasing, prognosis has improved significantly with 5-year survival outcomes improving from 40% to 90% over the past 5 decades.1 Breast cancer is a molecular diverse disease with at least 4 biologically distinct biological subtypes, each with individual clinical characteristics, therapeutic strategies, and prognoses.2 The St. Gallen Expert Consensus Panel advocate use of multigene expression assays [eg, OncotypeDX Recurrence Score (RS), Genomic Health Inc., Redwood City, CA] as the gold standard for molecular stratification,3 and immunohistochemical evaluation of estrogen (ER), progesterone (PgR), and human epidermal growth factor receptor-2 (HER2) receptors act as surrogate phenotypical biomarkers to decipher biological subtypes.4 Moreover, appraising receptor status is crucial for guiding therapeutic decision making.5,6

Breast cancer survival is equivalent for those treated with adjuvant and neoadjuvant chemotherapy (NAC).7,8 Nevertheless, advantages of NAC include potential tumor downstaging, increased patient eligibility for breast conservation surgery (BCS), improved resectability,9,10 and the generation of in vivo data regarding tumour sensitivity to systemic therapies, which carry prognostic significance.12,13 In particular, predicting response to NAC translates into important disease outcomes in the setting of HER2-positive (HER2+) and triple negative (HER2−) (15/120), and 20.8% triple negative disease (25/120). In total, 46.7% of patients responded to NAC (56/125) and 26.7% achieved a pathological complete response (pCR) (32/120). For patients with L/B and HER2, increased Let-7a predicted response to NAC (P = 0.049), while decreased miR-145 predicted response to NAC in HER2+ (P = 0.033). For patients with luminal breast cancers, reduced Let-7a predicted achieving a pCR (P = 0.037) and reduced miR-145 predicted achieving a pCR to NAC in HER2+ (P = 0.027).
molecular subtypes. Unfortunately, predicting response is challenging due to variability in host and tumour factors.

Micro ribonucleic acids (or miRNAs) are small (19–25 nucleotides in length), noncoding ribonucleic acid (RNA) molecules which modulate gene expression by post-transcriptional degradation or translational inhibition of messenger RNA. Aberrant expression of miRNAs regulate breast cancer development and miRNAs have the ability to maintain stability in several biological tissues. Moreover, miRNA profiling may be performed relatively simply and inexpensively using real-time quantitative reverse transcriptase polymerase chain reaction, supporting their suitability as clinical biomarkers.

Translational research efforts are focused on identifying novel biomarkers capable of predicting response to NAC to tailor treatment strategies to each patient’s needs. The Cancer Trials Ireland – Irish Clinical Oncology Research Group 10/11 (CTRIAL ICORG10/11) is a prospective, multicenter trial which recruited 120 patients treated with NAC for breast cancer. Expression levels of a predetermined miRNA panel were relatively quantified from bloods samples using relative quantification polymerase chain reaction at several predetermined timepoints during NAC.

The aims of this study were:

- To determine whether circulating miRNA expression levels successfully predict response to NAC.
- To establish whether subtype specific miRNA biomarkers predict response to NAC.

Methods

Study Design

This prospective, multicenter trial was overseen by Clinical Trials Ireland (CTRIAL ICORG10/11—NCT1722851), with recruitment at 8 treatment sites in the Republic of Ireland. Local ethical approval was granted in February 2008 (C.A.151) and in January 2014 (C.A.1012).

MiRNA expression profiles were measured from liquid biopsies taken at diagnosis (Timepoint 1, or T1) and halfway during NAC (Timepoint 2, or T2) (Fig. 1). Thereafter, the changes in miRNA expression levels between these timepoints were determined (T2 minus T1) and correlated with response to NAC.

Inclusion and Exclusion Criteria

Consecutive female patients aged 18 years or older indicated to undergo standard-of-care NAC for breast cancer were considered for inclusion. Patients had to be capable of providing informed written consent. Patients failing to meet these inclusion criteria were not considered for inclusion.

Histopathology and Molecular Subtyping

Breast cancer molecular subtypes were classified using the 12th St. Gallen Expert Consensus panel. In brief, tumors were evaluated for ER and PgR using immunohistochemistry as per the 2010 American Society of Clinical Oncology/College of American Pathologists (ASCO/CAP) consensus. HER2 status was identified by Herceptest (DAKO Agilent pathology solutions, Santa Clara, CA), with a score of 3+ considered positive. Any 2+ inconclusive results were confirmed using fluorescent in situ hybridization.

Ki-67 evaluation was performed in just a few cases using MIB1 antibody testing. Luminal disease [luminal breast cancers (LBC)] was classified as ER+/HER2−, Luminal/HER2+ disease (L/HER2) was classified as ER+/HER2+, HER2 disease (HER2+) was classified as ER−/PgR−/HER2+, and triple negative breast cancer (TNBC) was classified as ER−/PgR−/HER2−. American Joint Committee on Cancer (AJCC) version 8 was used for staging.

Response to Chemotherapy

Response to NAC was based on Miller-Payne classification. In addition, patients were classified as “responders” and “nonresponders”。“responders” were patients who achieved pathological complete response (pCR) or a >90% reduction in tumor cellularity on their specimen (Miller-Payne grades 4/5), while those with >90% were classified as “nonresponders” (Miller-Payne grades 1–3). Further subclassification was performed with respect to those who achieved a pCR (Miller-Payne grade 5) versus those who did not (Miller-Payne grades 1–4).

Venous Blood Sampling

Venous samples from the 120 patients were collected during a 3-year period (May 2011–April 2014). Whole blood liquid biopsies were collected at 2 independent timepoints:

- Timepoint 1 (T1) – at breast cancer diagnosis, before treatment with NAC.
- Timepoint 2 (T2) – halfway during NAC (Fig. 1).

Venous blood samples were collected in 3 mL ethylendiaminetetraacetic acid (EDTA) tubes and stored at the local Surgery Cancer Biobank. A study enrolment diagram illustrating recruitment is outlined in Supplementary Appendix 1. A (Supplemental Digital Content 1, http://links.lww.com/SLA/E85).

MiRNA Expression Panel

Following literature review, a 5 miRNA panel was selected for evaluation (Let-7a, miR-21, miR-145, miR-155, miR-191).

FIGURE 1. Scheme of timepoints at which venous sampling occurred during this study: Timepoint 1 (T1) which involved venous sampling at breast cancer diagnosis (and before standard-of-care with neoadjuvant chemotherapy), and Timepoint 2 (T2) at the halfway point during neoadjuvant chemotherapy.
and miR-195) based on their relevance in breast cancer at the time of trial design.\textsuperscript{20,21,33} Two additional miRNAs were used as validated endogenous controls (miR-16 and miR-345), with the intention of standardizing miRNA expression.\textsuperscript{24,34} Table 1 outlines the relevance of each miRNA selected for evaluation.

RNA Isolation and Storage

Total RNA was extracted from whole blood (1 mL) using Trizol (as per the manufacturer’s instructions). RNA concentrations were determined using spectrophotometry (NanoDrop ND-1000 Technologies Inc., Wilmington, DE), as previously described.\textsuperscript{20} RNA was then transferred to storage tubes, labelled, and stored at \( -70^\circ\text{C} \) in our Cancer Biobank.

Analysis of miRNA Expression Levels

For each sample, miRNAs were relatively quantified using quantitative reverse transcriptase polymerase chain reaction. TaqMan assays were used for the amplification of target miRNAs, as per manufacturer’s instructions (TaqMan Fast Universal Master Mix (2X), No AmpErase UNG: Applied biosystems, Foster City, CA).\textsuperscript{14,34} Assays were performed using AB7900HT (Applied Biosystems), as per the manufacturer’s instructions. Reactions were initiated with a 10-minute incubation at 95°C, followed by 40 cycles at 95°C for 15 seconds, and 60°C for 60 seconds. We utilised miR-26b as an inter assay control from a breast cancer cell line, which was included on each plate. All reactions were performed in triplicate. The threshold SD for intra-assay and inter-assay replicates was 0.3. The percentage polymerase chain reaction amplification efficiencies (\( E \)) for each assay were calculated using the slope of the semi-log regression plot of cycle threshold versus log input of cDNA (10-fold dilution series of 5 points), with the following equation, and a threshold of 10% above or below 100% efficiency was applied: \( E = (10^{-\text{slope}} - 1) \times 100 \). MiRNA expression levels were calibrated and normalized using endogenous controls, before expression levels were calculated using QbasePlus\textsuperscript{\textregistered} software (Biogazelle, Gent, Belgium) using the geNorm method. MiRNA analysis was performed blinded to clinical information.

Statistical Analysis

Data were analysed using R version 3.2.3. Differences in miRNA expression profiles between T2 and T1 were calculated.

### Table 1. The Relevance of the 5 Target-miRNA and 2 Endogenous Controls Included in our Predetermined miRNA Panel

| Target | References | MiRNA Function |
|--------|------------|----------------|
| Let-7a | Heneghan et al\textsuperscript{20,21} | Increased expression in treatment naive breast cancer patients versus controls and postresection |
| miR-21 | Heneghan et al\textsuperscript{20,21} | Known as a well described oncogenic miRNA |
| miR-195 | Heneghan et al\textsuperscript{20,21} | Increased expression levels in breast cancers relative to other malignancies and controls |
| miR-16 | McDermott et al\textsuperscript{33} | Endogenous control in human circulation |
| miR-425 | McDermott et al\textsuperscript{33} | Endogenous control in human circulation |

### Table 2. Clinicopathological and Surgical Data for all 120 Included Patients With Correlations With Response to Neoadjuvant Chemotherapy

| Parameter | Variable | Total | "Responders", n (%) | "Nonresponders", n (%) | \( P \) | pCR, n (%) | Non-pCR, n (%) | \( P \) |
|-----------|----------|-------|---------------------|------------------------|------|-----------|--------------|------|
| Total number | — | 120 (100.0) | 56 (46.7) | 64 (53.3) | — | 32 (26.7) | 88 (73.3) | — |
| Age (y) | Median (IQR) | 55 (48–63) | 55 (48.5–61.5) | 0.895 | 53 (47–63) | 55 (48–64) | 0.531 |
| Tumor size (mm) | Median (IQR) | 38 (28–54) | 39 (30–54) | 0.674 | 35 (25–48) | 40 (30–55) | 0.176 |
| Nodal involvement | — | 43 (35.8) | 21 (48.8) | 22 (51.2) | 0.919 | 16 (37.2) | 26 (60.5) | 0.069 |
| Positive | — | 76 (64.2) | 35 (46.1) | 41 (53.9) | 16 (21.5) | 61 (80.3) |
| Negative | — | 1 (0.8) | 0 (0.0) | 1 (100.0) | 0 (0.0) | 1 (100.0) |
| Missing | — | 64 (53.3) | 21 (32.8) | 43 (67.2) | 9 (14.1) | 55 (85.9) |
| Grade 1 | — | 57 (47.5) | 35 (61.4) | 22 (38.6) | 23 (40.4) | 34 (59.6) |
| Grade 2 | — | 63 (52.5) | 21 (33.3) | 42 (66.7) | 0.04* | 9 (14.3) | 54 (85.7) | 0.02* |
| Grade 3 | — | 67 (55.8) | 37 (55.2) | 30 (44.8) | 0.054 | 24 (35.8) | 43 (64.2) | 0.014* |
| Progesterone receptor | — | 55 (44.2) | 19 (35.8) | 34 (64.2) | 8 (15.1) | 45 (84.9) |
| HER2 receptor | — | 57 (47.5) | 35 (61.4) | 22 (38.6) | 23 (40.4) | 34 (59.6) |
| Molecular Subtype | — | 59 (49.2) | 16 (27.1) | 43 (72.9) | <0.001* | 6 (10.2) | 53 (89.8) | <0.001* |
| LBC | — | 21 (17.5) | 13 (61.9) | 8 (38.1) | 6 (28.6) | 15 (71.4) |
| L/H/HER2 | — | 15 (12.5) | 11 (73.2) | 4 (26.8) | 9 (60.0) | 6 (40.0) |
| HER2+ | — | 25 (20.8) | 16 (64.0) | 9 (36.0) | 11 (44.0) | 14 (56.0) |
| TNBC | — | 67 (55.8) | 37 (55.2) | 30 (44.8) | 0.054 | 24 (35.8) | 43 (64.2) | 0.014* |

*Denotes statistical significance.
The Shapiro-Wilks test was used to assess the distribution of data. Nonparametric analyses [i.e., Kruskal-Wallis test (used to compare medians among multiple groups) and the 2-sample Wilcoxon rank sum test (used for all 2-sample comparisons)] were used as appropriate to correlate miRNA expression with response to NAC. All analyses were 2-tailed and statistical significance was defined as $P < 0.050$.

RESULTS

Clinicopathological and Surgical Data

Overall, 120 patients were included. The median age at diagnosis was 55.0 [interquartile range (IQR): 48.0–63.0] and tumor size was 38.0 mm (IQR: 28.0–54.0 mm). Almost 50.0% of included patients had LBC (49.2%, 59/120), 20.8% had TNBC (25/120), 17.5% had L/HER2 (21/120), and 12.5% had HER2+ (15/120). Most patients underwent BCS (55.8%, 67/120) and axillary lymph node dissection (70.8%, 85/120). Table 2 illustrates clinicopathological and surgical data for included patients.

Response to NAC

Overall, 46.7% were “responders” to NAC (56/120). “Responders” were likely to have grade 3 ($P < 0.001$), ER− ($P = 0.002$), PgR− ($P = 0.004$), HER2+ ($P = 0.002$), and be L/HER2, HER2+ or TNBC molecular subtypes ($P < 0.001$) (Table 2).

In the overall cohort, no miRNAs correlated with response to NAC (Supplementary Appendix 1.B, Supplemental Digital Content 1, http://links.lww.com/SLA/E85). In L/HER2, increased Let-7a expression identified “responders” to NAC ($P = 0.049$). Similarly, decreased miR-21 expression trended toward significance for identifying “responders” in L/HER2 ($P = 0.064$). For HER2+, decreased miR-145 expression identified “responders” to NAC ($P = 0.033$) and decreased Let-7a expression trended toward significance for identifying “responders” ($P = 0.052$). Furthermore, increased miR-21 expression identified “responders” in TNBC ($P = 0.089$) (Fig. 2).

Pathological Complete Response to NAC

Overall, 26.7% of patients achieved a pCR (32/120). Those achieving a pCR were likely to be node negative ($P < 0.001$), grade 3 ($P < 0.001$), ER− ($P < 0.001$), PgR− ($P = 0.002$), HER2+
(P = 0.027), and be L/HER2, HER2+ or TNBC molecular subtypes (P < 0.001). Patients achieving a pCR were likely to undergo BCS (P = 0.014) (Table 2).

In the overall cohort, no miRNAs correlated with achieving a pCR (Supplementary Appendix 1.C, Supplemental Digital Content 1, http://links.lww.com/SLA/E85). In LBC, reduced Let-7a expression predicted achieving a pCR (P = 0.037), while increased miR-145 expression trended toward significance in predicting a pCR (P = 0.072). In HER2+, reduced miR-145 expression predicted achieving a pCR (P = 0.027), while reduced Let-7a expression trended toward significance in predicting a pCR (P = 0.062). In L/HER2, reduced miR-21 expression trended toward significance in predicting a pCR (P = 0.058) (Fig. 3).

DISCUSSION

Breast cancer's subclassification into distinct molecular subtypes is essential for personalizing treatment strategies.2,25 Once oncological control is established, efforts are focused on promoting personalized treatment. This represents the ideology of precision oncology and the aim of ICORG10/11 was to identify biomarkers of host response. Such a biomarker could allow individualization of therapy by identifying responders who should continue the course of treatment as planned, unlike their nonresponding counterparts who may benefit from halting NAC and proceeding to surgery. While Luminal A is consistently endocrine responsive,35 there was a traditional tendency to deliver chemotherapy to all patients with locally advanced and node-positive LBC.36 ICORG 10/11 pragmatically included LBC patients receiving NAC to elucidate responsible biomarkers. Unsurprisingly, pCR rates, the proportion of “responders,” and miRNA profiles which correlated with treatment response varied among molecular subgroups, therefore substratifying these subgroups based on their sensitivity to NAC, while laying the foundation for the discovery of novel miRNA-related molecular subtypes.37 Reduced miR-145 expression predicted patients who achieved a pCR to NAC in HER2+ (ER−/HER2+) disease (P = 0.027), coinciding with the results of the translational research NeoALTTO trial.38 In NeoALTTO, Di Cosimo and colleagues described increased plasma miR-145 expression of those achieving a pCR after 2 weeks treatment with Trastuzumab. Moreover, reduced miR-145 expression differentiated “responders” from “nonresponders” following NAC in this study (P = 0.033). Recent ASCO/CAP guidelines recommend NAC for patients with HER2+ cancers, with exceptions limited to those with T1a-b/N0 disease.39 Therefore, further validation

FIGURE 3. The difference in miRNA expression profiles between Timepoint 2 minus Timepoint 1 and the correlation with pathological complete response to neoadjuvant chemotherapy for each of the breast cancer molecular subtypes.

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of these preliminary results assessing the utility of miR-145 in delineating response to NAC is crucial before miR-145 may be used to guide therapeutic decision-making regarding NAC in clinical practice. Furthermore, such a biomarker may be useful in dividing HER2+ patients into two narrower molecular subtypes: Those likely to respond (and derive benefit from continuing NAC) and those unlikely to respond (and may be beneficiaries from early NAC cessation and progression to surgery). However, further interrogation of this biomarker in the next generation of translational research trials is mandated before fully establishing its value in predicting tumour sensitivity to NAC in HER2+ disease.

Increased expression of Let-7a segregated “responders” from “nonresponders” in L/H2R2 (ER+/HER2+) (P = 0.049). Interestingly, reduced Let-7a expression differentiated those achieving a pCR (P = 0.062) and “responders” in HER2+ (ER-/HER2+) disease (P = 0.052). While the current paradigm encourages the division of HER2+ breast cancers based on ER status, the use of Let-7a may provide further subdivision based on therapeutic response to NAC, as observed in ICORG 10/11 in correlating reduced miR-145 with the reduction in tumour volume in response to NAC. Furthermore, response to NAC predicts long-term oncological outcomes, therefore, it is unsurprising that Fuso et al demonstrated a survival advantage with increased Let-7a-5p expression in HER2+ disease (event-free survival: 61 vs. 36 mo, P = 0.050). Thus, this study supports the potential use of Let-7a profiling to inform response to NAC which may translate indirectly into long-term survival outcomes.

This study provides preliminary data regarding reduced miR-21 in L/H2R2 in predicting a pCR (P = 0.058) and aberrant expression of miR-21 in deciphering “responders” from “nonresponders” in L/H2R2 (P = 0.064) and TNBC (P = 0.089). This is a potentially anticipated finding: in previous ICORG10/11 data published from McGuire et al, miR-21 expression measured at T1 in isolation acted as an independent predictor of response to NAC (P = 0.036). Similarly in the Geparquinto trial, Müller et al reported increased baseline miR-21 expression differentiated “responders” from “nonresponders” in HER2+ disease, while Rodriguez-Martinez et al outlined the discriminative ability of miR-21 to predict pCR to NAC. Interestingly, upregulation of miR-21 via signaling of the MAPK pathway correlates with increased HER2 signaling, providing the rationale for changes in miR-21 expression to predict NAC response in HER2+ disease in ICORG10/11. Moreover, miR-21 is a promotor of proliferation, invasion and initiating epithelial-mesenchymal transition in TNBC, supporting the reduction in miR-21 correlating with “response” to NAC in TNBC.

In LBC, decreased Let-7a expression correlated with achieving a pCR (P = 0.037), while miR-195 expression failed to indicate sensitivity to NAC. These results refute the results from NeoALTO, where reduced miR-195 identified those who achieved a pCR to combined Trastuzumab and Lapatinib in HER2+ disease. Furthermore, mir-195 levels have correlated with lower grade, reduced proliferation, and “True” LBC. While McGuire et al described reduced miR-145 to correlate with “responders” in LBC, increased mir-145 expression trended toward significance for predicting pCR in LBC (P = 0.064). This emphasizes the importance of serial miRNA measurement to aid prediction of response to NAC, as demonstrated in the in vivo data from ICORG10/11 and NeoALTO. This study has several limitations: firstly, patients recruited were all habitants of a unique cultural European region, facilitating unavoidable selection from a small genetic pool, inherently limiting the translation of these results to a global level. Secondly, the predetermined miRNA panel evaluated included targets of most interest at the time of trial design. During the time elapsed between study initiation and completion, newer miRNA targets have been discovered. Thirdly, while this study provides insights into predicting response to NAC for each biomolecular subtype, this stratification of tumors into 4 subtypes was based on the 12th St. Gallen Expert Consensus. More recently, the 12th Expert Consensus Panel revised this subclassification to encompass 5 molecular subtypes. In addition, the seminal TAILORx and RxPONDER trials now facilitate RS to identify candidates with ER+/HER2− disease who may be spared systemic chemotherapies, whether in the neoadjuvant or adjuvant settings. Finally, although the goal of this study is to individualize NAC prescription based on circulating miRNA, there are of course preliminary results which will inevitably require validation before having actual influence upon therapeutic decision making in clinical practice.

The ICORG10/11 study is the first prospective, multicenter, neoadjuvant translational research trial conducted evaluating and illustrating the potential value of circulating miRNA measurement in predicting response to NAC in an Irish population. This study supports the ideology that circulating miRNAs may personalize therapeutic decision-making for patients indicated to receive standard-of-care NAC. The next generation of translational research trials should focus on interrogating the use of circulating miRNAs to inform tumor sensitivity to NAC, before adaptions to current therapeutic strategies may be made in clinical practice.

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up-wards or down-wards – so you cannot conclude that there is a
difference in median just because you reject the null hypothesis
(with a P-value less than your threshold). You might also reject
the null hypothesis because of a lack of independence, or because
distributions do not have the same shapes. As a preliminary
analysis, the shapes of the distributions should be investigated,
and if there are no significant differences in shape, then the
Kruskal-Wallis test might reveal the differences in medians. In
short, these aspects should be presented in the “statistical anal-
ysis” section.

Response from Matthew G. Davey (Galway, Ireland)

In response to the first part of your discussion, this trial
was designed over 10 years ago, when microRNAs were con-
sidered to be novel biomarkers. At the time of trial design,
perhaps 100 to 300 biomarkers had been previously described.
Currently, we know of 3000 to 4000 microRNAs. At the time
of the trial design, we pragmatically included the ones that were
most relevant to breast cancer pathogenicity, and these were the
ones we which perceived to be the most likely to inform ther-
apecic decision-making at that time. We are currently designing
the next phase of neoadjuvant translational breast cancer
research trials and have learned that predominating the bio-
markers of interest at the time of trial design may, in fact, not be
the best way to identify biomarkers for evaluation in such a trial.

With respect to the performance of the Kruskal-Wallis
analysis, the Department of Surgery has collaborated with our
local Department of Mathematics and Statistics, which was
responsible for performing the data analytics for the ICORG 10/
11 trial. Our colleagues in the Department of Mathematics and
Statistics have reassured us that they used the Shapiro-Wilk test
to determine whether microRNA expression data was in a
parametric or nonparametric distribution before non-parametric
data analytics were performed.

Bas Wijnhoven (Rotterdam, The Netherlands)

You also perhaps had the opportunity to get some tissue
samples. If you think that these microRNAs are shedding from
the tumor, it might be nice to have a look at the tissue before
neoadjuvant chemotherapy. Most of them had surgery; did you
get tissue biopsies from after the operation as well, and try to see
whether the differences make sense in the tissue and whether they
 correlated with the serum? What are the target genes of these
microRNAs, and can you explain, to some extent, the observa-
tions by looking at the target genes or whether the microRNAs
influence some tumor characteristics?

Response from Matthew G. Davey (Galway, Ireland)

That’s a very interesting point and it is something that we
have since considered. As I previously mentioned, we are now
designing the next generation of microRNA clinical trials to
evaluate the role in neoadjuvant chemotherapy decision-making.
We do intend to take a core biopsy at diagnosis, and then, at the
time of resections, as well as blood samples along the way. It’s
well established that microRNA expressions in the blood don’t
necessarily correlate with that in the tumor, which not only
speaks to the ubiquitous nature of microRNA expression but
also how non-specific their behavior may be. It will be very
exciting to compare the biopsies, from the initial diagnostic
biopsy to the one at resection, to see what the microRNA pro-
files are like and correlate them with the blood. What that will
inform, however, is yet to be seen. With respect to the targets of
these microRNAs, this analysis was unfortunately not captured
within the objectives or results of the ICORG 10/11 trial.

John Reynolds (Dublin, Ireland)

Regarding the association between microRNA 145 and
HER-2 positive tumors, is that a random hit or is there a bio-
logical paradigm that might underpin it?

Response from Matthew G. Davey (Galway, Ireland)

I don’t think that this result was a random hit for two
reasons. First, the NeoALTTO study, which is an even larger,
though similar, prospective neoadjuvant translational research
trial of 450 HER-2 positive patients, identified that differential
expression of miR-145-5p, which is of a similar structure to miR-
145, correlated with achieving a pathological complete response.
These findings correlate directly with the results we found in this
study. We understand miR-145 to be an oncomir, meaning it is a
pro-oncogenic biomarker. Taking that into account, as a tumor
is hopefully responding and downsizing to treatment, we would
then expect to see reduced miR-145 expression that correlates
with a reduction in the measurement of miR-145 in the circu-
ation. That would be a reasonable hypothesis as to why we have
observed these results in this study, and why this has been pre-
viously seen in another prospective clinical trial.