A Single Biopsy is Valid for Genetic Diagnosis of Eosinophilic Esophagitis Regardless of Tissue Preservation or Location in the Esophagus

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ABSTRACT

Background & Aims: A new gene expression profile test may distinguish eosinophilic esophagitis (EoE) and gastroesophageal reflux disease (GERD), but the optimal tissue preparation and biopsy location are unknown. We aimed to determine if formalin-fixed paraffin-embedded (FFPE) and RNA-later (RNAL) preserved specimens from newly diagnosed EoE patients have equivalent gene expression scores and whether scores vary by esophageal biopsy location.

Methods: We analyzed prospectively collected and banked esophageal biopsies from EoE patients and GERD controls. Paired FFPE and RNAL samples from the distal, mid, and proximal esophagus were used. RNA was extracted, and gene expression for a previously constructed 96 gene panel was quantified with a summary expression score. Scores were compared between EoE and GERD patients, between FFPE and RNAL samples, and between the different esophageal locations.

Results: A total of 72 samples, representing paired FFPE and RNAL specimens from 9 EoE cases and 3 GERD controls, were analyzed. Overall median gene expression scores were similar between FFPE and RNAL (238 vs 227; p=0.64), correlation was excellent between FFPE and RNAL (Spearman’s rho=0.90; p<0.001), and there were no differences by biopsy level. Median gene scores distinguished EoE from controls (134 vs 402; p=0.02), and overall agreement between preservation methods and EoE case status was perfect (kappa=1.0; p<0.001).

Conclusions: Gene expression scores were equivalent in FFPE and RNAL, and were also similar across three esophageal locations. This implies that a single biopsy in either FFPE or RNAL from anywhere in the esophagus may have the potential for genetic diagnosis of EoE.

Key words: Eosinophilic esophagitis – gene expression – RNA – paraffin – biopsy – transcriptome.

INTRODUCTION

Eosinophilic esophagitis (EoE) is a chronic immune-mediated clinicopathologic condition [1]. In order to diagnose EoE, current criteria (EoE) require symptoms of esophageal dysfunction and persistent esophageal eosinophilia (at least 15 eosinophils per high power field [eos/hpf]) after a high-dose proton pump inhibitor (PPI) trial, and with other potential causes of eosinophilia excluded [2, 3]. While these appear to be straightforward, in clinical practice the differentiation between EoE and GERD is difficult. There are no pathognomonic features of EoE, symptoms such as dysphagia, heartburn, and chest pain can both be present in both GERD and EoE, and even high levels of esophageal eosinophilia are not specific [1, 4-12]. Moreover, there is a complicated relationship between EoE and GERD, and both conditions can coexist in some patients [13].

Because of this difficulty, there has been extensive research interest in distinguishing the two conditions. To date, there has been examination of symptom scores [9, 14-16], tissue biomarkers [10, 17-24], and non-invasive biomarkers [25-
endoscopic findings, and final diagnoses were also recorded
samples per patient). From each esophageal level were selected for each patient (6
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by biopsy location.
that there would be no differences between the two tissue
in the proximal, mid, or distal esophagus. We hypothesized
whether gene expression scores would vary by biopsy location
specimens from newly diagnosed EoE patients would have
paraffin-embedded (FFPE) and RNA-later preserved (RNAL)
in the esophagus are unknown.
The aim of this study was to determine if formalin-fixed
paraffin-embedded (FFPE) and RNA-later preserved (RNAL)
specimens from newly diagnosed EoE patients would have
equivalent results on the gene expression profile panel, and
whether gene expression scores would vary by biopsy location
in the proximal, mid, or distal esophagus. We hypothesized
that there would be no differences between the two tissue
preservation methods, but that differences might be detected
by biopsy location.

METHODS

Study subjects and specimen collection
This was a case-control study analyzing biospecimens that
were prospectively obtained and stored in the University of
North Carolina (UNC) EoE Patient Registry and Biobank.
This resource was created and maintained during prospective
investigations of EoE from 2009-2014 [20, 31-33], where
subjects were enrolled if they had symptoms of dysphagia,
gastroesophageal reflux disease (GERD), or suspected EoE.
These studies were approved by the UNC IRB, and all study
subjects provided informed consent for participating prior to
undergoing endoscopy; this included consent for future use
of stored specimens. The present study analyzing the banked
specimens was also approved by the UNC IRB.
Patients with EoE were diagnosed as per consensus
guidelines [2, 3]. Specifically, they had to have symptoms of
esophageal dysfunction (dysphagia, food impaction,
heartburn, chest pain), esophageal biopsies with ≥ 15 eos/
hpf that persisted after a high-dose PPI trial (20-40 mg twice
daily of any of the available PPIs, prescribed at the discretion
of their clinician), and exclusion of other potential causes of
esophageal eosinophilia. GERD controls were patients who did
not meet criteria for EoE diagnosis, but who had heartburn- or
reflux-predominant symptoms.
At the time of the endoscopy, research protocol biopsies
from all subjects were obtained in order to bank tissue for
future use. We obtained specimens from the distal (3 cm
above the gastroesophageal junction [GEJ]), mid (10 cm above
the GEJ), and proximal (15 cm above the GEJ) esophagus
using standard large capacity forceps (RJ4; Boston Scientific,
Marlborough, MA). At each level, one biopsy fragment was
placed in formalin and subsequently embedded in paraffin,
and one was placed in RNA Later (RNAL; Life Technologies/
Thermo-Fisher Scientific, Grand Island, NY), frozen, and
stored at -80°C. For this study, paired FFPE and RNAL samples
from each esophageal level were selected for each patient (6
samples per patient).

In addition to tissue, patient demographics, symptoms,
endoscopic findings, and final diagnoses were also recorded
prospectively. Further, the esophageal eosinophil counts were
determined based on our previously validated methodology
[34]. The maximum eosinophil density (eos/mm²) was
quantified in five high power fields and then converted to an
eosinophil count (eos/hpf) based on a microscopy field size of
0.24mm², the most commonly reported size in the literature
[35].

RNA extraction and gene expression
For the FFPE tissue, 5 sections 10 microns thick were cut
for RNA extraction. For the RNAL samples, the entire biopsy
(approximately 8mm³) was used. Each sample was processed
with standard techniques to extract RNA. In brief, for RNAL
samples, the miRNeasy mini RNA extraction kit (Qiagen,
Valencia, CA) was used. The biopsies were transferred to a
microtube and macerated in a small volume of QIAzol Lysis
Reagent with a micropestle (Fisher Sci, Pittsburgh, PA). After
addition of chloroform, mixing, and centrifugation, the
aqueous phases were mixed with ethanol and then transferred
to a QIAcube (Qiagen) preloaded with RNeasy mini columns
for purification. For FFPE samples, samples were de-paraffined
with xylene and digested with Proteinase K and lysis buffer
(miRNeasy FFPE kit, Qiagen). The digested samples were then
passed thru gDNA eliminator columns and then transferred
to the QIAcube preloaded with RNeasy MinElute columns for
purification. RNA extraction from FFPE samples was 100%
successful provided that there was adequate tissue, defined
as the equivalent of a standard-sized biopsy specimen. RNA
concentration purity was measured by NanoDrop spectrometry
(TermoFisher) and then reverse transcribed into cDNA using
iScript cDNA Synthesis kit (Bio-Rad, Hercules, CA) as per the
manufacturer’s instructions.

Next, gene expression for a previously constructed 96
gene panel [29] was quantified (Eosinophilic Diagnostic
Panel [EDP], Diagnovus, Nashville, TN). Specifically, a set of
TaqMan probes for the 96 genes (including two housekeeping
genes, GAPDH and 18S rRNA), were pre-spotted on 384 well
fluidic cards (TaqMan Low Density Array Cards; TLDA, Life
Technologies, Foster City, CA). The qPCR was performed on a
ViiA7 cycler (Life Technologies) to determine gene expression
levels measured as Ct.
Finally, using this expression data, a summary score was
calculated by subtracting the housekeeping gene from the
Ct value of each gene of interest to acquire the ΔCt and
then summing their absolute values of the upregulated and
downregulated genes separately, as previously described [29].
A difference of the two sums was used to calculate the EDP
score with a score <333 diagnostic for EoE.

Statistical analysis
Descriptive statistics were used to summarize demographic,
endoscopic, and histologic characteristics. The median of
the maximum esophageal eosinophil counts was calculated both
overall and for each esophageal level. Median gene expression
scores were also calculated overall and for each esophageal
level, for both FFPE and RNAL preservation methods. The
overall score in a given individual was defined as the mean
of the scores from all levels. Using non-parametric methods
(Wilcoxon rank-sum; Wilcoxon sign-rank) the median gene

scores were compared between the EoE and GERD groups, between FFPE and RNAL samples, and between the different esophageal locations. We also assessed for differences in individual gene expression by the biopsy location and preservation method, requiring any differentially expressed genes to pass false discovery rate (FDR) [36]. Spearman’s correlation was performed between the FFPE and RNAL gene scores, as well as between the gene scores and esophageal eosinophil counts, both overall and by esophageal level. Finally, agreement between EoE case status (as defined by the consensus diagnostic guidelines [2, 3]) and gene score was determined using the kappa coefficient, both overall and by esophageal level. All analyses were performed with Stata 9.2 (StataCorp, College Station, TX).

RESULTS

Patients’ characteristics and samples
A total of 72 samples, representing paired FFPE and RNAL specimens from the proximal, mid, and distal esophagus from each of 9 EoE cases and 3 GERD controls, were analyzed for this study. Those with EoE were younger than GERD controls (median 34 vs 62 years; p = 0.03), all had dysphagia, and typical endoscopic findings such as rings, furrows, plaques, and edema were common (Table I). The median of the maximum eosinophil count was 80 eos/hpf in the EoE group and 0 eos/hpf in the controls (p < 0.001). For the controls, two subjects had normal biopsies with no esophageal eosinophilia, and one had a biopsy showing 6 eos/hpf. For the EoE cases, eosinophil counts were high at all esophageal levels (50, 60, and 49 eos/hpf for the proximal, mid, and distal esophagus, respectively).

Gene scores by preservation methods and biopsy location
When RNAL samples were compared to FFPE samples for the entire study population, there were no significant differences in the median gene scores either overall or by biopsy location (Fig. 1). For example, overall median gene scores were 227 for RNAL and 238 for FFPE (p = 0.64), and scores in the mid esophagus were 242 and 264, respectively (p = 0.43).

In addition, there were statistically significant correlations between the RNAL and FFPE scores, both overall and by biopsy location (Fig. 2). For example, Spearman’s rho was 0.90 for the overall correlation (p < 0.001), and 0.87 in the mid esophagus (p < 0.001). There were no differences in the expression of individual genes (none passed false detection rate) either by biopsy location or by tissue preservation type (gene array data are shown in the Supplemental Figure 1).

EoE diagnosis by preservation method and biopsy location
Median gene scores were significantly different between the EoE cases and the controls (Table I). For RNAL, EoE cases had a median score of 134 compared to 402 in the controls (p = 0.02), and for FFPE, the median scores were 223 and 362, respectively (p = 0.01). Additionally, there were no differences in these scores by preservation method among cases alone (p =

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* Medians were compared between groups with Wilcoxon rank-sum, and proportions were compared with Fisher’s exact test.

**No differences detected for comparisons between medial RNALater and FFPE scores for GERD controls (p = 0.29) or EoE cases (p = 0.21) using Wilcoxon signed-rank test.
or among controls alone (p = 0.29). Using a score <333 as the threshold for diagnosis of EoE, overall agreement between preservation methods and EoE case status was perfect, with a kappa of 1.0 (p < 0.001; Fig. 3). Agreement was also excellent to perfect at each esophageal level, with kappas ranging from 0.8 to 1.0 (p ≤ 0.01 for all comparisons).

There were also significant inverse correlations between the esophageal eosinophil count and the gene score for both RNAL and FFPE, both overall and at all esophageal levels (Fig. 4). Specifically, higher eosinophilic counts were associated with lower gene scores, and this association held regardless of case or control status. For example, there was one EoE subject with low proximal and mid esophageal eosinophil counts, but high distal counts (3, 6, and 110 eos/hpf, respectively). This subject had correspondingly higher gene scores at the proximal and mid- levels for both RNAL (468 and 426, respectively) and FFPE (358 and 330), but lower scores distally (262 for RNAL; 177 for FFPE).

![Fig. 1. Comparison of gene expression scores for samples preserved in FFPE (diamonds) vs RNALater (circles). (A) Overall score. (B) Score for the proximal esophageal biopsy. (C) Score for the mid esophageal biopsy. (D) Score for the distal esophageal biopsy. For all graphs, the solid black line represents the median value.](image)

![Fig. 2. Correlation of gene expression scores measured in FFPE and RNALater. (A) Overall score. (B) Score for the proximal esophageal biopsy. (C) Score for the mid esophageal biopsy. (D) Score for the distal esophageal biopsy. For all graphs, the dotted lines show the score cut point (333) below which a score is consistent with a diagnosis of EoE.](image)

![Fig. 3. Agreement between case status and tissue preservation method, as measured by kappa, for the overall gene score, as well as gene score by esophageal level. Dark gray bars show agreement between case status and RNALater, light gray bars show agreement between case status and FFPE, and black bars show agreement between RNALater and FFPE.](image)
**DISCUSSION**

The current diagnostic algorithm for EoE requires that both clinical and histologic features are present and that competing causes of esophageal eosinophilia are excluded [1-3]. However, it remains difficult to distinguish EoE and GERD clinically [4-12], and few techniques have been validated to do so [20]. The recent development of a gene expression profile test, however, holds great promise for doing just that [29], but whether it was optimal to use RNAL- or FFPE-preserved tissue, and whether there was a difference in expression by level of the esophagus, was not known. The goals of this study were to determine whether gene scores for FFPE specimens were equivalent to those from RNAL, and whether there was variability in gene expression scores based on biopsy location. The results were strong and consistent. First, RNAL and FFPE gene scores were not statistically different, either overall or by biopsy level, and the scores were highly correlated in individual patients, regardless of biopsy location. We also did not detect any differences in individual gene expression by level. Second, the gene scores almost perfectly distinguished the EoE cases from the controls. This has clinical implications for this methodology: one biopsy, regardless of the tissue preservation method or the esophageal location, can potentially be used to help diagnose EoE.

The genetic expression profile of patients with EoE, subsequently termed the EoE transcriptome, was first described by Blanchard and colleagues in a study of children with either active EoE, reflux esophagitis, or normal controls [30]. They demonstrated that there were approximately 340 upregulated and 230 downregulated genes characteristic of EoE. Additionally, using high-throughput whole transcriptome RNA sequencing techniques, Sherrill et al. recently expanded the genetic signature of EoE, identifying 1607 differentially expressed genes [37]. Identification of differential gene expression led to the development of a molecular diagnostic approach for EoE, recently published by Wen et al. [29]. In this landmark study, the technique was developed in 15 pediatric patients with active EoE and 14 normal controls, and then confirmed in an independent population of 18 EoE cases and 14 controls. Additional analysis showed similar results in 12 adults with EoE, and examined diagnostic utility in an additional 50 controls and 82 EoE cases. While the majority of samples assessed in this study were in RNAL or were fresh tissue, there were a subset tested in FFPE that yielded similar results. However, there was no assessment of expression at different levels of the esophagus. Our study assessed the same 96 gene panel in an adult population, using a similar but commercialized platform. As with the data from the Wen study, the panel was nearly perfect in its ability to discriminate...
EoE cases from controls, and we have expanded on their findings with our analysis throughout different levels of the esophagus.

This study has some potential limitations, as well as notable strengths, to address. Firstly, this was conducted at a single center and included only adults, so we are unable to comment on whether the same findings would be seen in other settings or in children. However, the study by Wen et al. presented comparable results for RNAL and FFPE in a pediatric population [29]. Our cases and controls are also not well-matched regarding age, and further validation will be needed in an age-matched population. Secondly, it is possible that the gene score is a marker of inflammation and may not be disease specific, but the study design does not allow us to comment on the specificity of the gene panel in non-EoE inflammatory conditions, including proton pump inhibitor-responsive esophageal eosinophilia. Thirdly, while the number of subjects included in this study was small, there were a large number of specimens analyzed and the results were robust and consistent across several analysis techniques. Moreover, this study utilized prospectively collected and banked tissue samples, with standardized protocols and storage methods, from well-characterized EoE cases and GERD controls. This is a clinically-relevant population in whom this gene panel could potentially be applicable, and allowed for a unique comparison between paired specimens from multiple esophageal locations with the two tissue preservation techniques. We were also able to perform analyses correlating gene scores with eosinophil counts, both overall and by esophageal level. We feel that these strengths outweigh the possible limitations of the study. Additionally, while EoE has been shown to be a patchy disease histologically, with wide variations in eosinophil counts throughout the esophagus [38], it appears that the gene expression may be more consistent. An assessment of gene expression related to variations in esophageal eosinophilia should be explored in future studies.

**CONCLUSION**

This analysis of prospectively collected and banked esophageal biopsy samples showed that gene expression scores in EoE cases and GERD controls were equivalent in FFPE and RNAL tissue. Further, gene expression scores were similar in the three esophageal locations tested, and correlated strongly with the esophageal eosinophil counts. This implies that a single biopsy in either FFPE or RNAL from anywhere in the esophagus may have the potential to be used for genetic diagnosis of EoE.

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**Authors’ contribution:** E.S.D.: project conception, study design, data/specimen collection, data analysis/interpretation, manuscript drafting, V.Y., M.A., and J.S.: specimen processing/analysis supervision; data analysis/interpretation. All authors: critical revision, final approval.

**Supplementary material** (Fig. 1) available from [http://www.jgld.ro/2015/2/8.html](http://www.jgld.ro/2015/2/8.html)

**REFERENCES**


Fig. 1. Gene expression read-outs by location and specimen type. Subjects are in columns, specific genes are in rows. Red colors indicate down-regulation (low expression) and yellow/blue colors indicate up-regulation (high expression)