INTRODUCTION

Ewing sarcoma (EwS) is a rare, aggressive sarcoma with a peak incidence in adolescents and young adults (AYAs). Primary tumors may arise in the soft tissue or bone, and staging differentiates patients with localized vs. metastatic disease. All patients receive multi-agent intensive chemotherapy and local control. Despite this treatment, over 20% of patients with localized disease and 60–80% of patients with metastatic disease suffer relapse and die of disease. In addition, those who survive are often left with debilitating late effects. Clinical features aside from stage have proven inadequate to meaningfully classify patients for risk-stratified therapy. Therefore, there is a critical need to develop approaches to risk stratify patients with EwS based on molecular features. Over the past decade, new technology has enabled the study of multiple molecular biomarkers in EwS. Preliminary evidence requiring validation supports copy number changes, and loss of function mutations in tumor suppressor genes as biomarkers of outcome in EwS. Initial studies of circulating tumor DNA demonstrated that diagnostic ctDNA burden and ctDNA clearance during induction are also associated with outcome. In addition, fusion partner should be a pre-requisite for enrollment on EwS clinical trials, and the fusion type and structure require further study to determine prognostic impact. These emerging biomarkers represent a new horizon in our understanding of disease risk and will enable future efforts to develop risk-adapted treatment.
To evaluate potential molecular prognostic biomarkers, we organized an international working group of disease experts to evaluate the most promising and clinically mature candidates in EwS and update a prior consensus published in 2013. Our working group reviewed the prognostic significance of a select group of contemporary molecular biomarkers with strong pre-clinical evidence including translocation subtype and mechanism of formation; STAG2 loss and TP53 pathogenic mutations; copy number variants; tumor mutational burden (TMB); circulating tumor DNA (ctDNA); and germline DNA defects (Fig. 1). We evaluated the current level of evidence for each available biomarker and identified molecular features that this expert panel agrees should be assessed in all future phase II/III studies. Biomarker data were evaluated in context of the relevant treatment regimen, with the understanding that 5-drug interval compressed therapy is now considered the preferred chemotherapy backbone in Europe and North America (Supplemental Table 1). These molecular biomarkers include those important for proper diagnosis and several that warrant prospective evaluation to validate prognostic value. This review is not intended to be comprehensive of all biomarkers in EwS, but rather focus on those judged to be most promising to inform risk-adapted therapy in the near future.

CURRENT DEFINITION OF EWS

Molecular diagnostics

The latest 2020 WHO Classification of Tumors of Soft Tissue and Bone includes four categories for round cell sarcomas: EwS; round cell sarcoma with EWSR1::non-ETS fusions; CIC-rearranged sarcomas; and sarcomas with BCOR gene alterations. Based on a recent international survey of pathologists and oncologists, most respondents defined EwS as harboring FET::ETS gene family fusions and agreed this group should define a primary therapy arm for a future frontline clinical trial. However, a consensus was not met on how to classify the remaining round cell sarcomas with molecular variants. This survey highlights the need for centralized molecular test standardization within clinical trials to provide a uniform approach in diagnosis, classification, and treatment of these patients.

Classic EwS is characterized by solid sheets of monomorphic round cells with ill-defined cell borders, scant, clear cytoplasm and round, uniform nuclei with open or fine chromatin. It typically lacks nuclear pleomorphism, spindling or epithelioid morphology. Immunohistochemically, most EwS tumors show diffuse and strong membranous CD99 positivity as well as NKL2.2 and FLI1 staining. The defining molecular feature of the tumor is the characteristic FET::ETS fusion involving: EWSR1::FLI1 (70–80%) and EWSR1/FUS:ERG fusions (15%), followed by EWSR1/FUS::FEV (5%), and EWSR1::ETV1/4 (1%)18. Ewing sarcoma (EwS) with ERG fusions may display a more variable spectrum of histologies, and due to the unbalanced translocation, FISH testing for EWSR1 gene rearrangements may show false negative results in more than half of the cases19. Such patients may require RNA or DNA sequencing approaches for fusion confirmation.

The incidence of sarcoma with EWSR1::non-ETS fusions accounted for 6% of the 240 patients in a recently analyzed large cohort of patients with round cell sarcomas with EWSR1 or FUS fusions18. Alternative EWSR1 rearrangements present diagnostic challenges with therapeutic implications and highlight the underperformance of EWSR1 break-apart FISH as a sole means for molecular diagnosis. For example, EWSR1::Patz1 rearranged tumors are epigenetically and genomically distinct from EwS with canonical fusions20,21. Although most tumors show positivity for CD99, a small subset of cases show focal or negative staining18. In addition, EWSR1::Patz1 sarcomas exhibit a divergent morphology with round and spindle cell features and a polyphenotypic immunoprofile which may pose significant diagnostic pitfalls and simulate other sarcoma types. EWSR1::N-FATc2 positive tumors are more common in older patients, characterized by monomorphic round to epithelioid cells in anastomosing cords and abundant myxohyaline to collagenous extracellular matrix. They harbor EWSR1 rearrangements and gains/amplifications which may serve as a diagnostic hint. The overall prognosis of these patients is poor18,27. CIC::DUX4 gene fusion, resulting from either t(4;19) or t(10;19) translocation, is the most common genetic abnormality detected in two-thirds of FET-negative round cell sarcomas. CIC-rearranged sarcomas occur most commonly in young adults and have an inferior outcome with a 5-year survival of only 43% vs. 77% for EwS (P = 0.002) in one series of 115 patients. Tumors have a predilection for soft tissue and show a variable round cell phenotype, admixed with epithelioid, spindle and myoid stroma components with immunopositivity for WT1 and ETV4 and variable CD99 expression. Molecular studies have highlighted the underperformance of FISH and RNAseq methods in diagnosing sarcomas with CIC gene abnormalities. Similarly, despite histological similarities, the BCOR family of tumors demonstrate distinctive clinical presentations and outcomes, including an overall favorable prognosis for patients with BCOR::CCNB3-positive tumors, and a highly aggressive behavior for BCOR ITD round cell sarcomas.
In summary, the presence of variant translocations may be difficult to detect or lead to the misclassification of tumors as EwS. Further, many variant translocations are not readily detectable by morphology or standard FISH. DNA and RNA-based sequencing approaches may be required in some cases for translocation identification/confirmation. Therefore, National Comprehensive Cancer Network (NCCN) guidelines recommend assessment of sarcoma pathology in a center with access to specialized testing, as these diagnoses are nuanced. Nevertheless, because the biology, outcomes and responsiveness to EwS-based treatment are distinct among patients with variant translocations, clinical trials in EwS should assess the primary objective in patients with FET::ETS fusion-positive EwS. Future phase II/III clinical studies must include molecular diagnostics for fusion partners to accurately classify patients as having EwS to allow the interpretation of trials designed for this tumor. Testing should aim to identify both fusion partners and preferably be performed at a centralized laboratory.

CLINICAL PROGNOSTIC FEATURES IN EWS

For patients with newly diagnosed EwS, stage remains the strongest prognostic factor. Patients presenting with metastatic disease have poor outcomes with a 2-year event-free survival (EFS) of 20–40%5,33. Among patients with metastatic disease, multiple studies have demonstrated that patients with isolated pulmonary metastatic disease have more favorable outcomes compared to patients with extrapulmonary metastasis34–36. In contrast, for patients treated on the most recent Children’s Oncology Group (COG) frontline trial for localized disease, 5-year EFS was 78%6.

Among the 70% of patients who present with localized tumors, the evidence for clinically useful prognostic factors is variable. In Europe, tumor size at presentation and histologic response have been used in clinical trials to differentiate patients with localized tumors into standard risk and high-risk groups57. Tumor size at presentation was prognostic in multiple studies58–60. Tumor volume of ≥200 mL was used on the recent European upfront trials to identify patients with high-risk localized disease. While prognostic on the most recent COG trial for patients with non-metastatic disease AEWS1031, patients with tumors ≥8 cm or ≥200 mL still had a 5-year EFS of 70% or greater and thus, in the context of the now international standard interval compressed 5-drug chemotherapy, tumor size holds modest prognostic impact61.

Response to therapy as assessed by histologic response following induction therapy is used to identify patients with high-risk localized disease in European trials61,62. However, in the context of 5-drug interval compressed chemotherapy, tumor necrosis was modestly prognostic with data demonstrating a 5-year EFS of 81% (95% CI 3–87%) and 75% (95% CI 4–81%) for patients with no viable tumor vs. any viable tumor at local control (P = 0.055)6. The evidence for radiologic response is variable63.

In addition, age at diagnosis, pelvic primary site, race, ethnicity, and sex have all been shown to carry modest prognostic significance64–66. These histologic studies are informative but cannot be fully generalized because these patients were treated with historic chemotherapy regimens, and eligibility relied upon histologic diagnosis and did not necessarily exclude patients with variant EwS translocations. Therefore, in the context of the current international standard of interval compressed therapy, these clinical risk factors provide only modest prognostic information and have not been used to identify patients with notably high or low-risk disease for testing risk-adapted therapy. Nevertheless, clinical features may be useful to identify low- and high-risk subgroups in patients with FET::ETS fusions when considered in the context of contemporary therapy and the molecular biomarkers described below.

SUMMARY OF EVIDENCE FOR MOLECULAR BIOMARKERS FOR RISK-STRATIFICATION AT DIAGNOSIS

Evaluation strategy for available biomarkers

The molecular biomarkers described in this review were evaluated by an expert working group from the North America and Europe. The relevant literature for each molecular biomarker was reviewed by the group who determined the strength of the available literature, viable assays through which the biomarker can be assessed, and whether the current state of evidence suggests the biomarker should be: (1) assessed only in the research space; (2) requires further investigation in a larger cohort; (3) requires prospective validation with a pre-defined statistical plan; or (4) can be used to define eligibility or disease subgroups for therapeutic stratification on a prospective trial. Given this was an international working group, strict use of biomarker language specific to regional regulatory bodies was avoided.

Translocation and chromoplexy

The FET-ETS fusion protein is the defining molecular feature of EwS (see molecular diagnosis section above). Most commonly, EWSR1::FLI1 forms from the t(11;22)(q24;12) chromosomal translocation joining exon 7 of EWSR1 to either exon 6 (type I; 60% of cases) or exon 5 (type II; 25% of cases) of FLI146. Alternative fusions occur between exons 7, 9, 10 of EWSR1 or exons 4–8 of FLI1, including a subset of tumors with a cryptic exon 8 (and intrinsic breakpoint) that is universally spliced out to yield a mature functional type I EWSR1::FLI146. In addition, alternative FET:ETS fusions have been described either involving the EWSR1 family member FUS or more commonly with other ETS family members such as ERG, ETVI, ETV4, and FEV47–52.

Early studies suggested that a type I EWSR1::FLI1 fusion might be prognostic and suggested a higher probability of relapse free survival for 31 patients with type I fusions relative to 24 patients with alternative fusions (RFS = 0.72 ± 0.1 vs. 0.21 ± 0.12; p = 0.04)55. A follow-up multivariate analysis of 99 patients with EwS supported this idea and reported a relative risk (RR) of 0.37 (P = 0.014) for type I fusions relative to other fusion types (RR = 0.32; p = 0.034)56. However, a larger data set of 119 patients treated with 5-drug chemotherapy in the COG failed to demonstrate a difference in either EFS or OS for type I vs. non-type I EWSR1::FLI1 fusions57. Consistent with this observation, the largest analysis to date of 565 patients treated on the Euro-EW.IL.N.G. 99 trial also failed to demonstrate a difference in relapse or death among patients with type I, type II or EWSR1::ERG fusions (P = 0.95 and P = 0.83)58. However, there was a slight, albeit non-significant increase in the risk of relapse or progression (HR, 1.38; 95% CI, 0.96–2.0 P = 0.1) or death (HR, 1.48; 95% CI, 0.98–2.2; P = 0.07) in 91 patients with EWSR1::FLI1 fusions that were called “EFx” and excluded type I, II and EWSR1::ERG fusions. This group was not captured in the COG analysis and includes the type III fusion protein (EWSR1 exon 10 fused to FLI1 exon 6). Importantly, the additional EWSRT exons lead to the inclusion of functional IQ and RGG domains that could influence sensitivity to either standard or targeted therapy59,60. Therefore, while a type I fusion is unlikely to be prognostic, the EFx subgroup (inclusive of type III fusions) remains an open question.

An intriguing possibility is that the mechanism of translocation could be prognostic. Although chromosomal translocations form by multiple mechanisms, in some tumors they result from catastrophic genomic events like chromosome shattering, called chromothripsis, or aberrant repair of loop-structures, known as chromoplexy59–61. In EwS, the occurrence of chromoplexy was found to be quite common, occurring in 42% of cases (52/124)62. This included all EWSR1::ERG rearrangements as well as a subset of canonical EWSR1::FLI1 fusions. Importantly, the presence of chromoplexy increased the likelihood of relapse (54% vs. 30%, p < 0.05)62. Consistent with the higher rate of relapse, there was
also an association with poor prognostic molecular features such as higher mutational burden and TP53 mutation (see TP53 section), but not STAG2 or CDKN2A mutations. Further, 504 differentially expressed genes distinguished EwS tumors formed by chromoplexy from “simple EwS” \( (P < 0.001) \), suggesting the activation of distinct biological pathways.

In summary, while the type I fusion is unlikely to be prognostic in the setting of interval compressed therapy, the type EFx subtype remains an open question. Further, processes that create fusions through catastrophic chromosomal events like chromoplexy lead to altered transcriptional profiles and may identify patients with inferior outcomes. Finally, as described above, variant translocations in tumors with inferior outcomes, such as EWSR1::NFIATc2 and CIC::DUX4 fusions, should be considered distinct biological entities and evaluated an unbalanced t(1;16) rearrangement. CNV studies within the landscape of these and other CNVs and their prognostic significance of chr 8 gains65,66,73. Some studies have suggested prognostic significance of chr 8 gains, including whole chromosome, segment 8q, MYC (8q24) and/or RAD21.66,67 Chr 8 gain66 and MYC amplification70 have been suggested to be more common in relapsed tumors. A study of 52 EwS specimens showed gain in chr 8 or chr 12 in 83% of relapsed specimens compared with primary tumor (47%) or metastatic specimens (42%) from initial diagnosis. In addition, chr 8 gain has been associated with a trend toward worse survival in two studies of 30 and 28 patients, demonstrating a non-significant decrease in 5-year EFS survival of 25–35% with chr 8 gain \( (P = 0.16) \).71,72 Studies from larger cohorts, however, have failed to demonstrate prognostic significance of chr 8 gains65,66,73.

In summary, CNVs are common in EwS and statistically significant differences in outcomes have been described, particularly for 1q gain and 16q loss. These prior studies are limited by patient numbers, and the lack of uniformity in the populations (localizes vs. metastatic) and treatment approaches. In addition, variations in testing methodologies challenges the comparison of the results across studies. These results highlight the need to further investigate the prognostic significance of CNV at these loci prospectively by a uniform approach to truly define the landscape of these and other CNVs and their prognostic implications in EwS. Multiple studies are in progress with the most notable being a prospective validation of 1q/16q using 1q and 16q specific FISH probes that is currently being performed using the EuroEwing 2012 cohort.

**Copy number variants 1q gain, 16q loss, 8q gain, other**

Several studies investigating copy number variations (CNV) in EwS identified recurring abnormalities involving whole chromosomes or segments including gains in chr 8 (50% of cases), chr 2 (25% of cases), chr 1q (25% of cases), and chr 20 (10–20% of cases)\(^{2,64}\). The most common deletion involves chr 9p and CDKN2A\(^2\). Chromosome 1q gain is frequently associated with 16q loss as the result of an unbalanced t(1;16) rearrangement\(^{64,65}\). CNV studies within the EuroEwing 99/2008 trial have shown that 1q gain and, possibly, 16q loss define patients with adverse outcomes\(^{64,66}\). 1q gain was detected in 55% of relapsed specimens compared to 11.5% of non-relapsed and was associated with inferior OS \( (P < 0.001) \); Fig. 2a\(^{64}\). A larger study verified this finding with 1q gain, demonstrating substantially inferior OS \( (P < 0.001) \); Fig. 2b\(^{65}\). Similarly, loss of 16q was associated with inferior OS (Fig. 2c; \( P = 0.0037 \)) and was co-associated with gain of 1q, although their combination did not show an additive effect on OS\(^{65}\). An attractive candidate on 1q is CDT2, a gene involved in cell cycle control whose gene dosage may increase proliferation rates in 1q gained EwS\(^{64,67}\). Survival curves demonstrating the prognostic impact of 1q gain and 16q loss are shown here adapted with permission from Tirode F. et al., Cancer Discovery, 2014\(^{65}\) (b). Survival curves for patient stratified by STAG2 and TP53 status, adapted with permission from Tirode F. et al, Cancer Discovery, 2014\(^{65}\) (c). Survival curves for patients stratified by STAG2 and TP53 status, adapted with permission from Tirode F. et al, Cancer Discovery, 2014\(^{65}\) (d).

**Fig. 2 Poor prognostic molecular features of Ewing sarcoma tumors.** Survival curves demonstrating the prognostic impact of 1q gain shown here are adapted with permission from Mackintosh C. et al., Oncogene, 2012\(^{64}\) (a) and Tirode F. et al., Cancer Discovery, 2014\(^{65}\) (b). Survival curves for 16q loss are shown here adapted with permission from Tirode F. et al., Cancer Discovery, 2014\(^{65}\) (c). Survival curves for patients stratified by STAG2 and TP53 status, adapted with permission from Tirode F. et al, Cancer Discovery, 2014\(^{65}\) (d).
patients with TP53 somatic nonsense and frameshift mutations in the EWSR1::FLI1 STAG2 expression in EwS alters the transcriptional program of the X chromosome were found to result in complete loss of whole genome, whole exome and other sequencing collected at diagnosis and relapse and in ctDNA, using Sanger, BJC Ewing sarcoma. Adapted with permission from Shulman DS. et al., BJ, 2018.

TP53 mutations in EwS
TP53 mutations have been identified in EwS tumor specimens collected at diagnosis and relapse and in ctDNA, using Sanger, whole genome, whole exome and panel sequencing methods. Detecting functional alterations in p53 by immunohistochemistry has been historically difficult because pathogenic mutations can result either in loss of protein expression when TP53 is deleted or truncated or high levels of nuclear protein accumulation in cancers with a missense mutation in TP53. Therefore, functional alterations of p53 are currently best detected by identifying pathogenic mutations by molecular laboratory methods.

Next-generation sequencing (NGS) studies of EwS cohorts demonstrated that pathogenic TP53 mutations are detected in 5–10% of cases. However, these studies included sample cohorts that were a mix of clinical phenotypes. Further studies are still needed to clearly delineate the rate of TP53 mutations in patients with newly diagnosed localized disease, metastatic disease, and at the time of relapse. Associations between TP53 mutations and outcomes in EwS have similarly been limited. Initial retrospective studies, including single-institutional studies, suggested an association of TP53 mutation with poor outcome. However, a more contemporary study conducted in collaboration with the COG was unable to identify a significant association between pathogenic TP53 mutations with outcome in patients with newly diagnosed localized EwS. This study demonstrated non-significantly inferior outcomes for the 8 patients with TP53 mutations (HR = 1.83 [95% CI: 0.65-5.19]) but was limited by cohort size (n = 96) and included sequencing of only TP53 exons 5 through 8.

Loss of STAG2 in EwS tumors
STAG2 loss occurs in 15–20% of primary EwS. Heterozygous somatic nonsense and frameshift mutations in the STAG2 gene on the X chromosome were found to result in complete loss of protein expression, presumably due to inactivation on the other X chromosome in female patients.

The STAG2 protein is a component of the cohesin complex, which has a role in chromosomal organization and segregation. Recent studies now demonstrate, as previously noted, that loss of STAG2 expression in EwS alters the transcriptional program of EWSR1::FLI1, resulting in a more invasive cellular phenotype.

Although STAG2 mutations are readily detectable by many existing NGS panels, loss of STAG2 protein expression has been documented in tumors that do not have any detectable mutations in this gene. The evaluation of STAG2 expression may be better assessed through standard immunohistochemical (IHC) staining of tumor biopsy samples. Indeed, the binary nature of expression within EwS cells makes the interpretation of IHC staining relatively straightforward, even allowing for the identification of areas of subclonal loss of STAG2 expression in biopsy samples. Furthermore, the ubiquitous nuclear expression of STAG2 in endothelial cells, present throughout viable EwS tumors, provides an internal positive control on every stained slide.

A previous study demonstrated that STAG2 mutation alone (n = 39) was associated with poor overall survival in a cohort of 299 patients with mixed clinical phenotypes (5-year OS ~50% vs. 70% for STAG2mut vs. STAG2WT; P = 0.007). Loss of STAG2 expression by IHC was enriched in patients with metastatic EwS in a retrospective single-institutional cohort of 59 patients with newly diagnosed disease. STAG2 mutations also appeared to be acquired or selected for at the time of relapse. Furthermore, by RNA-sequencing STAG2 loss of function gene signature correlated with poor outcome. A recent study of 108 patients with localized EwS previously treated on AEWS0031 demonstrated that STAG2 loss of expression occurred in 27% of patients, and 5-year EFS was 52% (95% CI 33–68%) and 75% (95% CI 63–84%) for patients with STAG2 loss vs. STAG2 expressed (P = 0.0018).

Loss of STAG2 and TP53 mutations
STAG2 and TP53 mutations co-occur more often than would be expected by chance. In one study of 299 patients with EwS, the combination of STAG2 loss and pathogenic TP53 mutations was associated with a worse outcome than patients with either single variant alone and relative to wild-type TP53 and STAG2. In one study, one patient was found to acquire different sets of STAG2 and TP53 mutations at different recurrences suggesting a clonal advantage for cells with both variants.

In summary, the clinical impact of STAG2 and TP53 mutations as potential prognostic biomarkers in EwS continues to rely on retrospective observational analysis. Prospective analyses of large clinically annotated cohorts in cooperative group studies will be required to fully test and validate their prognostic impact. Several ongoing efforts should provide the data needed to definitively determine the prognostic impact of STAG2 and TP53 in patients with localized EwS.

Circulating tumor DNA
Detection of cancer-derived ctDNA from cell-free DNA isolated from bodily fluids, such as plasma, CSF, and urine has been utilized in multiple cancer types. In EwS, the most common detection strategy quantitates the pathognomonic FET:EFS fusions by either PCR or hybrid capture targeted NGS. PCR-based assays designed to detect fusion breakpoint sequences are highly sensitive. However, the fusion breakpoints occur across broad intronic regions and are unique to each patient, requiring development of patient specific assays. Moreover, sequencing of tumor tissue is generally necessary for breakpoint identification. Targeted NGS panels of select EWSR1 introns obviate the need for a priori tumor profiling but are not as sensitive as PCR-based assays. In a series of studies evaluating the ability to detect ctDNA in patients with newly diagnosed localized and metastatic patients with EwS, ctDNA detection rates for PCR- and NGS-based assays were 137/146 (94%) and 61/100 (61%) of samples, respectively. In patients in whom ctDNA is detectable by both droplet digital PCR and targeted NGS assays, a strong correlation between the two methodologies has been demonstrated. More recently, integrated methods including genetic and epigenetic detection of ctDNA has been applied to

Published in partnership with The Hormel Institute, University of Minnesota
retrospective samples from patients with EwS, demonstrating highly sensitive and specific quantification of ctDNA. These methods show strong pre-clinical validation and require testing in larger clinical cohorts to determine clinical validity.

In one study of 50 patients with localized EwS, patients with detectable ctDNA by NGS (≥1.5%) had a 3-year EFS of 49% (95% CI: 24–69%) vs. 82% (95% CI: 49–93%) for patients without detectable ctDNA at diagnosis (Fig. 3). In a study of 102 patients with localized and metastatic EwS, ctDNA burden at diagnosis was divided into tertiles and higher ctDNA burden was associated with inferior outcomes. However, these ctDNA tiers were not prognostic among the 67 patients with localized disease in that study. Serial assessment of ctDNA demonstrated that patients who remained ctDNA positive at start of chemotherapy cycle 2 and 3 were more likely to have relapse events. Given the well-established ctDNA technology and strong preliminary data, prospective ctDNA-based biomarker studies should be incorporated into clinical trials for EwS as an integrated biomarker to confirm the prognostic relevance of specific ctDNA levels at diagnosis as well as the predictive value of ctDNA during treatment. Importantly, the translation of these research-based assays into a clinical laboratory environment will be needed if ctDNA analyses are to be utilized for risk-stratification. Given the long lead time required for clinical implementation of NGS and/or patient-specific MRD assay platforms, consideration should be given to starting this process now.

Tumor mutational burden

Ewing sarcoma (EwS) is characterized by a remarkably quiet genome, with a low TMB relative to most other malignancies. While recurrent mutations have been described in STAG2 and TP53,105,106 (see earlier), the overall mutational burden across the genome in these tumors is generally less than 1 mutation/Mb and nearly always <10 mutations/Mb. In this context, two studies have evaluated the potential prognostic impact of TMB in EwS. The first study utilized whole genome sequencing (WGS) to classify patients into tertiles defined by SNVs and indels, with a statistically significant association with overall survival. However, the effect size was relatively modest with 5-year overall survival estimates clustered within 20% points across all tertiles. Another group utilized WGS data and an analytic algorithm that provided an estimate of mutations that led to protein alterations. When analyzed in this way, patients with higher mutation burden had statistically significantly inferior overall survival, with a univariate hazard ratio of 2.6. Higher mutation burden was associated with older age and metastatic stage, though mutation burden remained prognostic on multivariate analysis.

Overall, the main limitations of utilizing TMB for clinical risk stratification are the paucity of data supporting this approach, the lack of substantial dynamic range in the marker, and the relatively modest effect size associated with this marker.

Germline DNA damage defects

Historically, unlike some other subtypes of pediatric sarcomas, EwS has not been associated with classic cancer predisposition syndromes. Exceedingly rare reports (n = 3) exist of siblings developing EwS.100,101 In one study, first-degree relatives of patients with EwS have an increased risk of developing cancers such as brain and female genital cancers, and second-degree relatives demonstrated an increased risk of breast cancer among others.102 Genome-wide association studies have identified genetic haplotypes associated with an increased risk of EwS, which possibly contribute to disease onset through generating higher affinity binding sites for EWSR1::FLI1 as shown at CIC::DUX4 cis-regulatory elements

In the past ~5 years, an emerging subset of patients with EwS and pathogenic germline variants in genes involved in DNA damage repair have been noted, due in large part to multiple large-scale sequencing efforts in the pediatric oncology population and the increase in patient tumors being sent for sequencing upon relapse.107–114 Examples of genes impacted in this patient cohort include but are not limited to FANCC, FANCM, BRCA1, BRCA2, RAD51, BARD1, SLX4, and PALB2. While any individual gene variant is rare, as a group, these pathogenic germline variants are found in ~10–13% of patients with EwS.

Clinically, this subset of patients is of high interest given the possibility that these patients may respond differently to therapy as compared to patients without an additional deficit in DNA damage repair. Intriguing future questions to address include: (1) Are overall patient outcomes different? (2) Would this cohort be more likely to respond to certain DNA damaging agents/combinations in the setting of relapse? (3) Does this cohort experience more toxicities from treatment? (4) If outcomes are better in this cohort, should a study of therapy reduction be considered?

When considering the future of risk-stratifying patients with EwS, patients with pathogenic germline variants in DNA damage repair genes are a logical group of patients to study prospectively on future clinical trials.

FUTURE DIRECTIONS

Justification for tissue collection

These studies together highlight molecular features that may stratify patients into higher and lower risk therapeutic treatment cohorts. The limitations of most published studies include sample size and challenges in integrating data across studies due to the non-homogeneous mixture of patient treatment protocols and variations in the molecular approach to analysis. Centralized tissue collection and analysis would mitigate many of these issues. Indeed, centralized translocation testing is justified based on the nuanced approach to diagnosis described above and the need to exclude patients with alternative fusions such as CIC::DUX4, or cases with EWSR1::non-ETS fusions that would appear as EwS if only a break-apart FISH is used. Other promising biomarkers such as STAG2 and TP53 require state of the art IHC and genomic analyses to identify all the various pathogenic mutations. Further, the detection of copy number changes and the quantitation of ctDNA can each be performed with multiple techniques. Therefore, uniformity in the analysis is required before statistically valid and biologically meaningful conclusions can be drawn about any of these biomarkers. A centralized approach would provide high-quality, outcome-linked, standardized tissue collection that will facilitate further pre-clinical analysis of these and other molecular features as technologies and science evolve. The hope is that this will drive discovery of more effective and less toxic precision therapy.

Inclusion of centers large and small

Marked disparities in access to, and quality of care contribute to differences in prognosis and outcome for patients who are afflicted with serious illnesses, including cancer.115 If not recognized and mitigated, these disparities can be especially profound for patients who suffer from rare diseases. Given their relative rarity, sarcomas, including EwS, present diagnostic challenges and centralized pathology review has been shown to greatly increase diagnostic accuracy in several cooperative sarcoma studies.116–118 In the US, the NCCN guidelines now recommend diagnosis at an institute with access to NGS testing. In addition, patients with EwS receive complex multi-disciplinary care that most often requires travel to an urban center, a factor that amplifies inequities. Moreover, given the unique challenges faced by AYA patients, the increased prevalence of EwS in
As our knowledge of fundamental biologic processes and targeted therapeutics grows, the critical role of advanced molecular tools for the diagnosis and treatment stratification of sarcomas, including EwS, becomes increasingly important. As detailed in this review, there are many new and innovative biomarkers that may, with further study, prove useful for prognostication in newly diagnosed and relapsed patients. If so, this will enable the development of new clinical trials and treatment strategies that could finally improve survival for patients with high-risk disease and lessen unnecessary treatment-related toxicity for low-risk patients. To ensure that all patients with EwS benefit from these advances, it will be essential that access to these innovative diagnostic and prognostic tools be available to all patients. The centralized nature of pediatric cancer care in most high-income countries, alleviates some disparities, though the need to travel to large urban centers remains a major inequity. Therefore, it is an ethical imperative that the future testing, validation, and implementation of prognostic and predictive biomarkers be achieved through an equity lens. As such, we propose that the need for equal access to centralized pathologic review and tumor testing should be considered as essential elements of any new assay or approach.

Clinical trial integration
A refined risk-stratification approach to EwS inclusive of biologically relevant molecular features would have substantial implications for clinical trial design, especially in the localized patient population. There exists sufficient prognostic information and readily available assays to justify prospective evaluation of multiple biomarkers, as pre-specified trial aims, in all patients to define biologically relevant subtypes. As the above molecular biomarkers undergo further study, one could consider how they might be incorporated into a future risk-adapted frontline clinical trial should these biomarkers be successfully validated. Here, we propose a hypothetical schema based on available data that incorporates both clinical and molecular features. The schema utilizes three risk groups: a

![Fig. 4 Hypothetical risk-stratified treatment schema.](image_url)

We provide a hypothetical risk-stratification schema that one could envision as the biomarkers described in this manuscript are validated. This schema incorporates clinical and molecular biomarkers.
low-risk, standard-risk and high-risk group. The low-risk group would include patients with an estimated 5-year EFS of ≥90%. Reasonable approaches to de-escalation would include reduction of alkylators, doxorubicin or etoposide to reduce fertility, cardiac, and second malignant neoplasm-related late effects. The standard risk-group would include patients with an estimated 5-year EFS of 50–90% who should continue to receive current standard risk chemotherapy (i.e., interval compressed VDC/IE) with or without additional trial interventions of minimal risk, such as maintenance therapy. The high-risk group would include patients with a <50% 5-year EFS and receive the standard chemotheraphy backbone with the addition of a targeted agent and/or novel approaches to extending or intensifying chemotherapy (Fig. 4). Beyond prognostication, the promise of serial biomarker assessments and response-based interventions including changes in ctDNA, and tumor volume, paired with examination of tumor tissue viable populations after induction chemotherapy should be incorporated into trials.\(^{78,80,94,122}\). Given that most patients with EwS go into a radiographic remission prior to disease relapse, these efforts must prioritize understanding molecular minimal residual disease. Collection of specimens from patients with disease relapse must also be prioritized to improve our understanding of mechanisms of resistance. Translational teams should determine the best, non-consumptive, correlative studies to evaluate mechanisms of resistance and relapse to advance our understanding of tumor evolution and acquired resistance with epigenetic changes from STAG2 in particular.\(^{87,88,90,123}\)

**CONCLUSIONS AND PERSPECTIVE**

In summary, we have assembled an international working group of disease experts to review available molecular biomarkers with the potential to be used in combination with clinical factors to identify disease subgroups. Many of the molecular biomarkers covered in this review have strong preliminary evidence as prognostic biomarkers in EwS. Translocation was the only biomarker with strong enough evidence to warrant use as an integral biomarker for enrollment to future clinical trials for the purposes of identifying patients with EwS, separate from EWSR1 round cell sarcoma with non-ETS fusion partners, CIC::DUX4 sarcomas and BCOR::CCNB3 sarcomas. Translocation type and chromoplexy remains an area of active investigation and requires further study before being used for risk-stratification. Co-occurrence of 1p:\(^{12}\) and 16q:\(^{12}\) have strong preliminary evidence and are currently being validated on the EuroEwing 2012 trial prospectively. Similarly, STAG2 is the most prognostically-valuable single gene candidate and is being evaluated in EuroEwing 2012 trial and in a large cohort of previously treated patients from AEWS1031 and AEWS0031, alone and when co-mutated with TP53. ctDNA is the most promising peripheral blood biomarker, potentially representing a measure of micrometastatic disease, with preliminary evidence from two large cohorts from the United States and Europe. Prospective validation of diagnostic ctDNA burden is underway. We note that while our review was focused, many other promising prognostic biomarkers beyond this review have been proposed, including but not limited to SOX2, mir-34a, Ki67, neurexin-1, RRM2, PRC1, IGFI/IGFBP3, and may also warrant further evaluation.\(^{24–131}\)

These biomarkers, primarily studied over the prior decade, represent promising markers for testing of risk-adapted treatment approaches but require definitive validation prior to use for assigning therapeutic strategies. These studies will provide the first step in the realization of risk-adapted treatment strategies. Therefore, our review highlights the need for comprehensive evaluation of these biomarkers in large, annotated cohorts with pre-planned analysis such that the prognostic impact of each marker can be definitively determined in the context of relevant clinical features. Such efforts are ongoing and will inform future attempts to implement risk-adapted therapy. The ultimate success of these strategies will be greatly enhanced through collaborative science and international harmonization of approaches to biomarker implementation.

Received: 1 March 2022; Accepted: 19 August 2022; Published online: 17 September 2022

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ACKNOWLEDGEMENTS
The authors would like to acknowledge Dr. Federico Comitani for contribution of the circo plot in Fig. 1.

AUTHOR CONTRIBUTIONS
All authors contributed equally to this work. All authors participated in study development, interactive teleconferences, writing and editing. Overall study design was performed by D.S.S., D.R., E.R.J., K.A.J., and P.J.G. Final paper assembly and editing was done by D.S.S. and P.J.G.

COMPETING INTERESTS
The authors declare no competing interests.

ADDITIONAL INFORMATION
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41698-022-00307-2.

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