Atypical teratoid rhabdoid tumours (ATRTs) are the most common malignant central nervous system tumours in children ≤1 year of age and represent approximately 1–2% of all pediatric brain tumours. ATRT is a primarily monogenic disease characterized by the bi-allelic loss of the SMARCB1 gene, which encodes the hSNF5 subunit of the SWI/SNF chromatin remodeling complex. Though conventional dose chemotherapy is not effective in most ATRT patients, high dose chemotherapy with autologous stem cell transplant, radiotherapy and/or intrathecal chemotherapy all show significant potential to improve patient survival. Recent epigenetic and transcriptional studies highlight three subgroups of ATRT, each with distinct clinical and molecular characteristics with corresponding therapeutic sensitivities, including epigenetic targeting, and inhibition of tyrosine kinases or growth/lineage specific pathways.

**Key Words :** Brain neoplasms · Rhabdoid tumour · SMARCB1 protein, Human · Antineoplastic agents · Epigenomics · Protein-tyrosine kinases.

**INTRODUCTION**

Atypical teratoid rhabdoid tumours (ATRTs) are malignant embryonal tumours of the central nervous system (CNS) characterized by bi-allelic loss of function alterations of SMARCB1, which encodes the hSNF5/BAF47/INI1 subunit of the SWI/SNF chromatin remodeling complex, and more rarely SMARCA4, which encodes the SWI/SNF subunit BRG1. ATRTs represent one of the most common and aggressive brain tumours of early childhood[25]. First identified as a distinct tumour type in 1996[49], ATRTs were difficult to distinguish from other embryonal brain tumours (EBTs) due to similar neuroepithelial histology. Despite ATRT’s designation as a separate disease entity in the World Health Organization’s Classification of CNS tumours[23, 42] since 2000, until recently, there has been limited progress in biological understanding and clinical management of this highly lethal tumor. Recent global transcriptional and methylation profiling of large tumor cohorts by two groups revealed ATRT comprises molecular subclasses with distinct clinicopathologic features, providing a much awaited breakthrough in clinical and biological understanding of ATRTs[32, 58].

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Elizabeth Anne Richardson, B.Sc,1,2 Ben Ho, M.Sc,2,3 Annie Huang, M.D., Ph.D.1-5

Department of Laboratory Medicine and Pathobiology,1 Faculty of Medicine, University of Toronto, Toronto, Canada

Arthur and Sonia Labatt Brain Tumour Research Centre,1 Hospital for Sick Children, Toronto, Canada

Department of Cell Biology,3 Hospital for Sick Children, Toronto, Canada

Department of Paediatrics,4 University of Toronto, Toronto, Canada

Division of Hematology/Oncology,5 Hospital for Sick Children, Toronto, Canada
CLINICAL PRESENTATION

It is difficult to accurately estimate incidence of ATRT in part due to its rarity and misdiagnosis as other EBTs; however recent advances in awareness and availability of molecular or histological tests allow for more accurate identification of ATRT. These tumours represent anywhere from 1–2% of all pediatric CNS tumours\(^{17}\), comprising up to 6.1% of malignant high grade tumours\(^{68}\). Overall, there is a slight male predominance of nearly 2 : 1\(^{10}\), though this does not carry to the subgroup level\(^{50}\). ATRT is primarily a disease of infants and toddlers, with a median age of diagnosis between 1.2 and 2.3 years\(^{13,46,64,68}\) and roughly 2/3 of all patients are diagnosed at \(\leq 3\) years of age\(^{57,68}\). ATRTs have been identified throughout the CNS, with supratentorial presentations being most common and spinal presentations the most rare at 1–7%\(^{2,57}\). Classic ATRT morphology is diverse, with epithelial, mesenchymal and neuroepithelial features. "Rhabdoid cells" (hallmark of nearly all rhabdoid tumours), may account for only a very small percentage of the cells observed in ATRTs, with some tumours exhibiting mostly small round blue cell morphology similar to those seen in medulloblastoma (MB). As a result ATRTs were often diagnosed as what was then known as PNET or as MB prior to the establishment and introduction of SMARCB1 as a molecular marker\(^{6}\).

Currently accepted diagnostic requirements of ATRT are genetic bi-allelic loss and/or negative immunohistochemistry staining of SMARCB1 or SMARCA4 and their respective gene products hSNF5/INI1/BAF47 and BRG1\(^{2,25}\). Approximately 20–35% of ATRT patients are found to carry bi-allelic germline alterations of SMARCB1 or SMARCA4; children with consequent “rhabdoid tumour predisposition syndrome” have a propensity to develop additional intra- and extra-cranial rhabdoid tumours at a young age\(^{5,8,14,27,37,51}\).

TREATMENT AND PROGNOSIS

There is presently no standardized treatment regimen for ATRTs. Retrospective data highlight the highly malignant nature of this disease with observed survival estimates at one year of \(\leq 50%\)\(^{13,64}\). With increased use of hSNF5 immunohistochemistry as a diagnostic tool and recognition of the need for more aggressive therapy for ATRTs, survival of correctly diagnosed ATRT patients after multimodal therapies including surgery, chemotherapy and radiotherapy has improved\(^{(p=0.047)\(^{68}\). However much remains unknown about the prognostic role of specific patient features or treatments\(^{40}\).

Currently available clinical literature on ATRTs is almost entirely retrospective, with frequently small and heterogeneously treated cohorts, from which only age, tumour location and metastatic status at diagnosis have been identified as key prognostic factors. Age \(>3\) years at diagnosis correlates with better prognosis\(^{3,28,29,57}\), possibly due to sparing of radiotherapy as well as reported increased incidence of metastasis in younger patients\(^{10,13,28,57}\). Furthermore, patients with rhabdoid tumour predisposition syndrome have been reported to carry a poorer prognosis. The extent to which age is a prognostic factor is still debated, with some groups reporting 50% disease free survival in younger patients\(^{11,67}\), while another identified age \(<2\) years as a prognostic factor only when combined with M-stage and Claudin 6 immunopositivity\(^{40}\). Favourable outcomes have also be linked with supratentorial tumour location\(^{10,13,46,64}\), while poorer outcomes are reported in the 14–21% of patients presenting with metastases at diagnosis\(^{2,9,28,57}\).

Extent of surgery is widely reported as a major prognostic indicator in ATRT\(^{28,29,37,68,69}\), with gross-total resection (GTR) frequently correlated with improved outcomes compared to subtotal (STR), near total (NTR), or partial resections (PR)\(^{10,48}\). In one study, GTR and PR patients exhibited a median event free survival (EFS) of 14 months and 9.25 months respectively\(^{20}\). Some studies, however, do not report prognostic impact from extent of surgery\(^{28,30}\), while Dufour et al.\(^{13}\) noted surgical extent was only a prognostic marker in univariate but not multi-variate analysis. These discrepancies highlight the gap in collective prospective data to evaluate the importance of clinical and treatment variables to patient prognosis.

Conventional dose chemotherapy

Conventional dose chemotherapy has been largely non-curative for ATRT patients in first-generation studies including the North American CCG9921 and POG 9923 trials which only observed a 10% EFS\(^{71}\). A substantially better 1-year progression-free survival of 53% was reported with use of a sarcoma-based regimen by the Dana-Farber group which included doxorubicin and dactinomycin in a “modified IRS-III” protocol\(^{10}\). While some groups have reported benefits of methotrexate and anthracycline based protocols\(^{90,95,96}\), others have noted no
survival differences. Similarly conflicting reports also exist regarding use of platinum and alkylating agent regimens in ATRTs. There is no consensus on the most promising and active agents for ATRTs, in part due to the heterogeneous, multi-agent therapies frequently administered to ATRT patients, but mostly because large scale clinical trials to robustly examine the relative contribution of these multiple variables have not been possible in this rare disease.

**High-dose chemotherapy (HDCT)**

HDCT with autologous stem cell rescue (ASCR) was initially adopted to defer cranio-spinal irradiation in patients <3 years, and has become an increasingly popular mainstay of ATRT treatment. High dose methotrexate has been included in some variations in the induction phase of treatment, including the Head Start (HS) II and HS II* studies, followed by consolidation therapy of 1–3 cycles of HDCT with carboplatin, etoposide and thiotepa. Patients enrolled in the earlier HS I regimen (6/6 dead of disease) had worse outcomes than their HDCT HS II counterparts (3/7 alive, no evidence of disease [NED]), notably with long term radiation free survival. Two major registry studies have examined impact of HDCT, with the first noting 46% of patients survived with NED, amongst survivors 50% had GTR and 33% received radiation. In the second, a Canadian registry study, HDCT conferred a survival benefit with 2-year overall survival improved from 27.3±9.5% to 47.9±12.1% compared to conventional chemotherapy treated patients (p=0.036). Although the benefit of HDCT may be difficult to separate from other favorable factors such as GTR and M0 status in 5/9 and 6/9 respectively, it is important to note that both studies demonstrate radiation free survival in a disease classically thought to be incurable. Based on these initial observations, the North American Children’s Oncology group conducted a HDCT prospective ATRT trial; preliminary reports from this ACNS0333 study is promising, with significantly improved survival observed in comparison to CCG9921 and POG9923 studies which used conventional chemotherapy approaches. Tekautz et al. also reported improved survival in children >3 years treated at St. Jude’s Children’s Research Hospital with 4 tandem HDCT regimens and cranio-spinal radiation when compared to historical cohorts. Superior survival with high dose chemotherapy in ATRTs has also been recently reported by Sung et al., who note that increased intensity of HDCT may allow for reduced craniospinal radiation in older children without compromising survival.

**Radiation therapy (RT)**

RT is typically deferred, avoided or dose and/or volume reduced in treatment of younger patients with ATRTs in order to avoid the associated neurocognitive toxicity; some protocols have also applied RT in a risk adapted manner. The role of RT in ATRT therapy is hotly debated as mixed results have been reported in national and institutional series. In the Canadian cohort reported by Lafay-Cousin, 6/11 patients were long term survivors after HDCT treatment without any RT. Similarly a German study showed no demonstrable survival benefit of RT on survival, von Hoff et al. also reported no significant difference in patients who received focal versus craniospinal irradiation (n=10, n=19, p=0.578), nor between those who received upfront versus salvage RT (p=0.314), thus suggesting that a group of ATRT patients may not need RT at all. However, Athale et al. reported in a meta-analysis that there was a trend towards greater mean survival time (18.4 months vs. 9.5 months, p=0.097) with RT. Buscariollo et al. similarly reported survival benefit for ATRT patients given RT (p=0.02). Thus, the benefit, dose and volume of RT in ATRT, and the choice of patients in which RT is needed remain unresolved.

**Intrathecal (IT) chemotherapy**

IT chemotherapy has also been explored both for prophylaxis and treatment of metastatic disease as an alternative to radiation of the CNS axis. Various combinations or single IT agents including methotrexate, cytarabine and/or hydrocortisone have been incorporated into several regimens, including the Dana-Farber ATRT protocol and intra-ventricular methotrexate in the German HIT SKK protocol. Contradictory findings are reported across studies, with a meta-analysis suggesting IT chemotherapy confers survival benefit while other individual studies suggest no additional benefit. However, as RT and/or HDCT are often applied together with IT chemotherapy in many of these regimens, the contribution of IT therapy alone to outcomes is difficult to discern. Interestingly, an institutional radiation-free treatment protocol at the Hospital for Sick Children in Canada, which uses monoagent IT during induction and in a prolonged maintenance – mirroring the practice in leukemia, has yielded surprising and encouraging survival in patients with M0 or M+ disease without use of RT.
evasive in prospective studies, this strategy to spare cranio-
spinal radiation would represent a significant step towards re-
ducing toxicity for ATRT survivors.

**MOLECULAR CHARACTERIZATION**

Initial genetic screens by Rorke et al.\(^\text{49}\) identified monosomy 22 in rhabdoid tumours in 1996, leading to the discovery of *SMARCB1* as a major tumour suppressor and etiologic gene in rhabdoid tumours including ATRT\(^\text{60}\). Previous studies suggested that loss of function of hSNF5 does not lead to genomic instability in cancer cell lines\(^\text{40}\), which is consistent with recent next generation sequencing studies which show ATRT genomes are highly stable with a very low coding mutation rate\(^\text{40}\). The clinical heterogeneity seen in ATRTs despite its monogenic etiology emphasizes the extent to which epigenetic dysregulation resulting from loss of SWI/SNF components drives this devastating disease.

Recent high resolution molecular studies have begun to reconcile observed clinical heterogeneity with the relatively bland genome of ATRTs by noting substantial heterogeneity in transcriptional and epigenetic profiles. In 2011, Birks et al.\(^\text{71}\) identified a subgroup of ATRT with high expression of bone morphological protein (BMP) pathway genes that correlated with shorter survival times. Torchia et al.\(^\text{80}\) performed a much larger scale integrated analysis of clinical and transcriptional data from 259 patients and noted two major transcriptional groups of ATRT – one primarily supra-tentorial group characterized by neurogenic differentiation and high ASCL1 protein expression, and a second primarily infra-tentorial group with enriched BMP signatures, which they termed group 1 and group 2 ATRTs respectively. Johann et al.\(^\text{32}\) and Torchia et al.\(^\text{80}\) subsequently independently reported on genetic, epigenetic and transcriptional characterization of more than 300 patients which revealed ATRTs could be subdivided into three methylation subgroups, with distinct clinical characteristics and targetable pathways (Fig. 1). Patient age distribution within each subtype is similar in both studies, with the lowest median patient age in ATRT-tyrosinase (TYR)/group 2A ATRTs while ATRT-MYC/group 2B ATRTs have the broadest patient age range and the most patients above 3 years of age. Neither group noted any subgroup predominance for metastasis or gender, as previously noted. Both the “ATRT-sonic hedgehog (SHH)” and “group 1” ATRTs have primarily neurogenic profiles, with increased Notch expression seen in both cohorts, which may indicate a potential therapeutic target for this subgroup. Though a comprehensive analysis comparing the two studies are pending, the ATRT-SHH, ATRT-TYR and ATRT-MYC subgroups seem to correspond with the gene enrichment signatures seen in group 1, group 2A, and group 2B subgroups respectively\(^\text{32,58}\). These high resolution genetic analyses did not reveal any additional recurrent coding alterations, however differences in both global and *SMARCB1* genotypes across ATRT subtypes were noted. ATRT-SHH/group 1 ATRTs demonstrated a higher frequency of focal alterations on *SMARCB1*, whereas in contrast the ATRT-MYC/group 2B subgroup harbors broader deletions on chr22q encompassing large portions of *SMARCB1* and surrounding genes.

A recent study by Han et al.\(^\text{30}\) created the first transgenic mouse model of ATRT by exploring the temporal deletion/inactivation of *Smarcb1*. They observed that *Smarcb1* knockout at E6–10 resulted mainly in intra-cranial tumours resembling ATRTs, with occasional extracranial tumours reminiscent of malignant rhabdoid tumours (MRTs)\(^\text{25}\). This study reinforces that the epigenetic mechanism associated with hSNF5 loss drives ATRT formation and additionally suggests different targeted cell of origin may contribute to the heterogenous nature of ATRTs. Further studies in *Drosophila* melanogaster have identified increased expression of upstream regulators of the Hippo signalling pathway when the fly homolog for *SMARCB1* (*snr1*) was knocked down, denoting an additional potential therapeutic target\(^\text{31}\). Experimental studies have also shown that *SMARCB1* loss leads to de-regulated expression of enhancer of zeste homolog 2 (EZH2), a histone methyl transferase which is a critical component of the PRC2 complex that antagonizes SWI/SNF activity during normal development\(^\text{67}\). Importantly, double knockouts of *SMARCB1* and *EZH2* induced senescence *in vitro* in MRT cell lines and prevented tumour formation in mice, indicating EZH2 as a promising therapeutic target.

**ATRT-BIOLOGY TARGETED THERAPEUTICS**

Improvement in ATRT outcomes has come with substantial toxicity associated with intense chemotherapy and/or radiation, therefore further therapy escalation may be difficult using conventional agents. Despite intensification of various modalities, up to 50% of ATRT patients have early disease progression and
Fig. 1. Summary of clinical, epigenetic and therapeutic sensitivity of ATRT subgroups. ATRT-SHH: sonic hedgehog subgroup, ATRT-TYR: tyrosinase subgroup, ATRT-MYC: MYC subgroup.
underscore the heterogeneous tumor biology inherent in this disease. Thus, further improvement in ATRT survival and functional outcome of survivors will need to come from incorporating and combining novel biologies with conventional treatments. To date, a spectrum of epigenetic and signalling inhibitors have been identified based on studies in ATRT or MRT cell lines - many of which are under evaluation in clinical trials.

Epigenetic inhibitors

Given that rhabdoid tumours are almost exclusively an epigenetically driven disease, targeting epigenetic regulatory mechanisms has been of great interest in ATRT and MRT treatment. Previous studies have shown that EZH2 expression is increased and required for tumour progression in hSNF5 deficient tumours\(^\text{67}\), and further that EZH2 inhibitors such as EPZ-6438 and 3-deazaneplanocin A prevent proliferation either alone, or in combination with other chemotherapeutics respectively\(^\text{35,61}\). EZH2 inhibitor tazemetostat is currently under evaluation in phase 1 clinical trials for a spectrum of SWI/SNF defective tumors including MRTs and ATRTs. Bromo/BET domain inhibitors such as JQ1 has also been shown to attenuate rhabdoid tumour growth via the Hedgehog pathway\(^\text{33,56}\), while other studies have noted success using histone deacetylase inhibitors (HDACi) such as trichostatin A, SAHA and SNDX-275\(^\text{41}\). Torchia et al.\(^\text{58}\) investigated a panel of small molecule drug and drug-like epigenetic inhibitors in three group 1 and five group 2 cell lines and observed a distinct sensitivity of group 1 cell lines to UNC999, UNC0638, and JQ1, while the HDACi LAQ824 reduced proliferation across all eight tested cell lines. Thus, it is likely that specific classes of epigenetic drugs may be more effective against one molecular class of ATRTs versus another.

Targeted and multi-kinase inhibitors

Observations that SMARCB1 loss leads to de-repression of Cyclin D1\(^\text{40}\), and lack of tumor formation in Ccnd1\(^/-\). Smarcb1\(^/-\) first demonstrated Cyclin D1 was essential for rhabdoid tumour growth presented Cyclin D1 as a promising therapeutic target\(^\text{60}\). These observations were further validated by observation of decreased Cyclin D1 expression and inhibition of rhabdoid tumor growth with combined treatment of the broad CDK inhibitors flavopiridol and tamoxifen\(^\text{33,54}\). These studies have led to evaluation of CDK4/6 inhibitor ribociclib in MRTs with some success in clinical studies\(^\text{20}\) and plans for evaluation in an upcoming prospective North American Children’s Oncology Group consortia clinical trial for ATRT. Pharmacologic inhibitors of the mitotic serine/threonine kinase Aurora A (also regulated by SWI/SNF, hSNF5) has also been of therapeutic interest\(^\text{40}\). In preclinical studies conducted by the NCI Pediatric Preclinical Testing Program, alisertib (MLN8237), an Aurora A kinase inhibitor exhibited IC50s <100 nM in BT12 and CHLA-266 ATRT cell lines and sensitized these cells to radiation\(^\text{62}\). Preliminary studies in patients with recurrent disease have been promising\(^\text{66}\), which has led to an ongoing phase 2 trial evaluating alisertib in recurrent ATRT and MRT at St. Jude’s Children’s Research Hospital.

Due to the highly heterogeneous nature of ATRTs, multi-tyrosine kinase inhibitors (multi-TKIs) are attractive for potentially reducing escape mechanisms or resistance. Imatinib, a first generation multi-TKI, was reported to inhibit cell growth via c-abl in G401 and A204 rhabdoid lines\(^\text{60}\). Similarly in vitro studies of ATRT cell lines have identified dual vascular endothelial growth factor/dual specificity mitogen-activated protein kinase kinase 1 (VEGF/MEK) inhibitors sorafenib and sunitinib as promising agents when tested alone or in combination with irinotecan, a topoisomerase I inhibitor\(^\text{30}\). Torchia et al.\(^\text{58}\) recently reported that second generation multi-TKIs nilotinib and dasatinib reduced cellular proliferation at nanomolar concentrations specifically in group 2 ATRTs via inhibition of platelet derived growth factor receptor beta (PDGFRβ), which is differentially epigenetically regulated across subgroups. These observations suggest that tumor sub-group context will be important to consider in evaluation of targeted agents. The identification of TKIs with well characterized safety profiles, previous use in pediatric oncology, as well as demonstrated blood brain barrier permeability suggest they are attractive candidates for rapid translation into clinical management of ATRTs.

Growth and lineage specific pathways

Interest in targeting growth and lineage signalling pathways has come from initial observations of an insulin-like growth factor (IGF) autocrine-paracrine loop in ATRTs between IGF-IR and IGF-2\(^\text{45}\), and more recent studies by Torchia et al.\(^\text{58}\) which showed ATRT subtype specific cell lines were sensitive to inhibition of NOTCH and BMP signalling. Specifically a role for IGF-IR in ATRT therapies has been supported by knockdown or inhibition of IGF-IR by antisense oligonucleotide as well as treatment with IGF-IR inhibitor NVP-AEW541 which induced death and decreased proliferation in BT12 and BT16.
ATRT cell lines\textsuperscript{1,2,3}.

Torchia et al.\textsuperscript{58} tested the functional significance of subtype specific enrichment of NOTCH and BMP signaling in group 1 and 2 ATRTs and observed that \(\gamma\)-secretase inhibitor DAPT selectively inhibited growth of group 1 ATRT cell lines and reduced expression of NOTCH pathway markers NICD, HES1, and HES5 in a dose-dependent manner. Further, they observed that dorsomorphin, a selective BMP pathway inhibitor, reduced cellular proliferation and expression of BMP markers BAMBI, SOST, and pSMAD1/5 in group 2 cell lines only\textsuperscript{58}. These experiments highlight future possibilities of exploiting lineage specific growth dependencies for ATRT therapies.

**FUTURE DIRECTIONS**

The last two decades have seen incredible advances in the understanding of ATRT biology and promising improvements in treatment of the disease. However, overall survival for ATRT is modest even with current approaches of maximal tolerated dose escalation. With greater understanding of subgroups of ATRTs and their molecular characteristics, more precise, biology-driven therapies can be developed to more effectively treat ATRT patients without unnecessary toxicity.

Identification of molecular classes of ATRTs with different therapeutic susceptibility represents a first step towards refinement in therapy through development and use of drugs matched to group specific biology. Classification of well-established ATRT cell lines into molecular subgroups\textsuperscript{58} and progress in development of \textit{in vivo} models of ATRT\textsuperscript{25,26} will continue to critically inform and guide testing of potentially translatable therapies. Stratification in future trials will require not only matching potential therapeutics with sensitive ATRT subtypes but including prognostic impact of subtype with other clinical risk factors. Whether specific ATRT subgroups can be treated with less aggressive therapies and without radiation, and which subgroups require further development of biology-specific treatments remain important, outstanding questions that will require large scale clinical trials. Global collaborative efforts to conduct large prospective cohort trials with robust subtyping and evaluation of prognostic impact will be critical to fully realize the therapeutic potential of risk stratified, biology-tailored ATRT therapies.

**CONFLICTS OF INTEREST**

No potential conflict of interest relevant to this article was reported.

**INFORMED CONSENT**

This type of study does not require informed consent.

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