Etoposide, an anticancer drug involved in therapy-related secondary leukemia: Enzymes at play

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Etoposide is a semi-synthetic glycoside derivative of podophyllotoxin, also known as VP-16. It is a widely used anticancer medicine in clinics. Unfortunately, high doses or long-term etoposide treatment can induce therapy-related leukemia. The mechanism by which etoposide induces secondary hematopoietic malignancies is still unclear. In this article, we review the potential mechanisms of etoposide induced therapy-related leukemia. Etoposide related leukemogenesis is known to depend on reactive oxidative metabolites of etoposide, notably etoposide quinone, which interacts with cellular proteins such as topoisomerases II (TOP2), CREB-binding protein (CREBBP), and T-Cell Protein Tyrosine Phosphatase (TCPTP). CYP3A4 and CYP3A5 metabolize etoposide to etoposide catechol, which readily oxidizes to etoposide quinone. As a poison of TOP2 enzymes, etoposide and its metabolites induce DNA double-stranded breaks (DSB), and the accumulation of DSB triggers cell apoptosis. If the cell survives, the DSB gives rise to the likelihood of faulty DNA repair events. The gene translocation could occur in mixed-lineage leukemia (MLL) gene, which is well-known in leukemogenesis. Recently, studies have revealed that etoposide metabolites, especially etoposide quinone, can covalently bind to cysteines residues of CREBBP and TCPTP enzymes. This leads to enzyme inhibition and further affects histone acetylation and phosphorylation of the JAK-STAT pathway, thus putatively altering the proliferation and differentiation of hematopoietic stem cells (HSC). In brief, current studies suggest that etoposide and its metabolites contribute to etoposide therapy-related leukemia through TOP2 mediated DSB and impairs specific enzyme activity, such as CREBBP and TCPTP.

Etoposide is a commonly used anticancer drug

Etoposide is a fundamental and essential part of combination chemotherapies for treating numerous cancers, such as lung cancer, lymphoma, leukemia [1]. Like most anti-neoplastic drugs, etoposide treatment has limited single-agent activity [2,3]. It is mainly associated with cisplatin, carboplatin, and cyclophosphamide in combination chemotherapy [4-6]. Currently, 4 to 6 cycles etoposide and cisplatin combination is the standard therapy commonly used for most small cell lung cancer in the clinic, with a 50%-80% objective response rate [7,8]. In combination with several drugs (lomustine, methotrexate, and prednisone), etoposide is also proposed to be first-line therapy in patients with non-Hodgkin’s lymphoma with no major cardiotoxicity [9,10]. Similarly, in Hodgkin’s disease, etoposide is a first-line chemotherapeutic agent in combination with other chemotherapeutic agents (vincristine, chloramphenicol, and prednisolone), was positive (77% response rate) [11]. In addition, it is reported that etoposide was active against gestational trophoblastic disease [12]. Etoposide was used for breast cancer treatment, the single-agent trial in untreated patients showed a response rate of approximately 15% in untreated patients [13]. Oral etoposide has been investigated in many clinical trials for the treatment of ovarian cancer response rates have varied in different studies (20.4% to about 30%), and it seems that the activity of this drug depends to a large extent on the degree of prior treatment [14].

Although etoposide has been widely and successfully used to treat many types of cancer, patients treated with etoposide may develop secondary leukemia. Due to the increases in the overall cure rate of patients, interest has arisen on the adverse effects, and special attention

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has been focused on the potential risk of therapy-related secondary leukemia [15–17] and the FDA (U.S. Food and Drug Administration) and EMA (European Medicines Agency) recommend different doses and treatment periods for different cancers as well [18, 19].

Metabolism of etoposide in human body

In cells, the oxidation of etoposide mainly involves cytochrome P450 3A family enzymes and/or peroxidases, such as myeloperoxidase (MPO). CYP3A4 and CYP3A5 are members of the cytochrome P450 superfamily of enzymes (encoded by the CYP3A4 and CYP3A5 genes). CYP3A4 and CYP3A5 metabolize etoposide to etoposide catechol, which is readily oxidized to etoposide quinone by cellular peroxidases [20]. The induction of CYP3A4 and CYP3A5 can enhance the clearance rate of etoposide [21, 22]. Etoposide catechol is oxidized by MPO in liver or bone marrow lysosomes [23, 24]. Etoposide can be directly metabolized to etoposide quinone with the assistance of prostaglandin-endoperoxidases 1/2 (PGTS1/2) or by other peroxidases (such as MPO) [25]. Oxidative metabolites of etoposide have a more potent inhibitory activity on DNA TOP2 cleavage near the MLL translocation breakpoint and more remarkable oxidizing ability than etoposide [26]. Etoposide can be cleared by UGT1A1 (UDP-glucuronosyltransferase family 1 member A1) etoposide glucuronides. In addition, etoposide quinone may be transferred to glutathione via GSTT1/GSTP1 (glutathione S-transferase). These two conjugated metabolites seem to inactivate the biological properties of etoposide [27, 28]. (Fig. 1)

Mechanism of action of etoposide

Early in the development of etoposide, its mechanism of action was considered similar to its parent compound, podophyllotoxin. These compounds inhibit microtubule assembly by preventing tubulin polymerization and then destroying spindle fibers. When the cells are exposed to podophyllotoxin, sister chromatids cannot be separated during mitosis because of a missing spindle [29]. As a result, cell division is arrested in mitosis with an increased cell number in metaphase. Nevertheless, cells exposed to etoposide show a decreased cell number in metaphase rather than an increase [30]. Simultaneously, studies reported that a low concentration of etoposide blocks the cell cycle in the late S or early G2 phase [31, 32]. These are significant indications that etoposide has a different mechanism compared to podophyllotoxin.

Currently, it is well known that etoposide is a TOP2 inhibitor, this inhibition being considered as a major anti-cancer mechanism of etoposide. TOP2 (alpha and beta isoforms) are ubiquitous enzymes that play a vital role in many basic DNA processes and regulates the under- and over-winding of DNA during DNA replication. TOP2 inhibitors play an integral role in cancer treatment due to the collapse/collision of the resultant bifurcation and unsolved DSB, leading to cancer cell apoptosis. Generally, TOP2 enzymes create a transient TOP2-DNA cleavage complex (TOP2cc) and resesals TOP2cc rapidly. Once the inhibitor interacts with TOP2, the TOP2cc is trapped, leading to an accumulation of DSB. If the DSB cannot be repaired timely, the cell will initiate apoptosis and eventually die [33, 34]. In contrast, if the DSB cell survives, it gives rise to the prospect of faulty DNA repair events, and the gene translocation could occur in the MLL gene, which is known for leukemogenesis. Chromosomal translocation causes abnormal growth and leads to the development of malignant tumors and neoplasms. More than 10,000 gene fusions have been identified in cancer [35, 36]. Etoposide and other chemotherapeutic do not directly cause translocation. However, the DSB caused by these molecules must be repaired, and it is the aberrant repair that can result in a translocation. (Fig. 2)

Topoisomerase II and topoisomerase II poisons

Topoisomerases are classified according to whether they cut one (type I) or two strands (type II) of DNA. Human cells express two TOP2 subtypes, TOP2α and TOP2β. The two subtypes share 70% amino acid sequence identity and encoded by two different genes. TOP2α is encoded by TOP2A, which is located at chromosome 17. TOP2β is encoded by

![Fig. 1. The main metabolic pathway of etoposide in vivo.](image-url)
TOP2B, which is located on chromosome 13 [37–39], TOP2α is highly expressed in proliferating cells and can be used as a biomarker of cell proliferation. While, TOP2α is poorly expressed in quiescent and differentiated cell populations [37]. In contrast, TOP2β expression was uniformly distributed in all cells [37,40]. Therefore, TOP2α is considered an effective target of TOP2 inhibitors, its impairment being considered as a contributor to the anticancer effects of etoposide.

TOP2 enzymes cleave DNA double-strands, which may cause DNA double-strand breaks [41,42]. Chemicals that inhibit TOP2 activity and decreases the levels of TOP2cc are termed TOP2 catalytic inhibitors. Chemicals that increase the levels of TOP2cc can convert the enzyme into a potential "toxin" that further produces DNA double-strand breaks are called TOP2 poisons [43].

TOP2 poisons can be grouped into two classes based on their mechanism of action. Interfacial poisons, such as etoposide, doxorubicin, or mitoxantrone bind non-covalently to the TOP2cc at the interface of protein-DNA complex and prevent TOP2 from rejoining the DNA ends [44,45]. Covalent poisons such as certain quinones (BQ or etoposide quinone) react with TOP2 at cysteine residues (albeit reaction with other amino acids cannot be ruled out) [46]. Covalent poisons produce irreversible TOP2cc and increase the levels of DSB by altering the conformation of TOP2 [47].

All clinically used TOP2 targeted anticancer drugs impair activities of both TOP2α and TOP2β [37]. However, the extent to which any TOP2 poison targets TOP2α and TOP2β and the relative contribution of either isoform to the drug’s therapeutic effect is not well understood [46,47]. It is worth noting that cellular and in vivo studies indicate that TOP2β is the primary enzyme responsible for gene breaks in MLL (see below).

Etoposide induced therapy-related leukemia

Leukemia is a carcinoma of the human hematopoietic system, comprising the bone marrow and lymphatic system. The precise causes of leukemia are not understood; cellular disorders and environmental factors are certainly involved [48]. Therapy-related leukemia has been under investigation for decades and is known to be induced by radiation and/or chemotherapy [49]. Drugs that induce secondary leukemia include alkylating agents, TOP2 poisons (for instance, etoposide and teniposide), cyclophosphamide, anthracyclines, and doxorubicin [15]. Etoposide is a well-known TOP2 poison and a commonly used anticancer drug. It is also recognized that etoposide can be leukemogenic, notably through MLL gene translocation [26,50].

Etoposide was approved for clinical use by the FDA in 1983, and a first report published in 1987 showed that etoposide treatment has a risk of induction of secondary leukemia in 1987 [51]. Subsequently, studies were published on etoposide and secondary leukemia, particularly acute myeloid leukemia (AML). Le Deley’s study showed that a high concentration and continuous etoposide treatment of solid tumors give a higher risk of treatment-related acute myeloid leukemia (t-AML) [52]. The study performed by N. J. Winnick et al. revealed that within 23 to 68 months after etoposide treatment, 10 out of 205 children developed secondary AML, and there was a 5.9% ± 3.2% risk of developing leukemia in the next four years [53]. Sugita et al. reported a high incidence of etoposide-associated secondary leukemia in children with non-Hodgkin’s lymphoma treated by etoposide. In order to reduce the associated secondary AML, the authors recommend giving etoposide treatment twice a week [54]. Kollmannsberger et al. demonstrated that patients receiving etoposide doses in excess of 2 g/m² had a 1.3% likelihood of developing s-AML [55]. Meanwhile, Ratain et al. proposed that high-dose etoposide is leukemogenesis in non-small cell lung cancer treatment, and the median dose of etoposide is 6795 mg/m² [51]. A study by M.A. Smith et al. found that the cumulative risk of secondary leukemia in six years was observed 3.3% in the low etoposide group, 0.7% in the medium etoposide group, and 2.2% in the high etoposide group, respectively (low: < 1.5 g/m²; Medium: 1.5 to 2.99 g/m²; high ≥3.0 g/m²), indicating that factors other than cumulative dose seem to be the main determinants of secondary leukemia risk [56]. Interestingly, it was revealed that receiving etoposide at doses greater than 4000 mg/m² tended to increase secondary acute non-lymphoblastic leukemia (s-ANLL). Meanwhile, all leukemias described in the Italian Langerhans cell histiocytosis (LCH) group are acute promyelocytic leukemia, which are not identified in Austria, Germany, Netherlands, and Switzerland (AGDS) groups [57]. It is reported that secondary leukemia does occur after conventional doses of etoposide treatment, but the low incidence does not change the risk-benefit ratio of etoposide-based chemotherapy in germ cell carcinoma [58]. Based on these researches, there is evidence that high-dose and long-term doses are critical factors in secondary leukemia caused by etoposide treatment, while other factors cannot be ruled out.
Topoisomerase II-associated DNA double-strand breakage and repair mechanisms

A critical intermediate in topoisomerase activity is the cleavage complex. Once topoisomerases have cleaved the DNA, each subunit of the TOP2 becomes covalently attached to the broken end of the 5′-phosphate group [40]. The cleavage complex is usually transient and naturally results from circumstances that are not clear or induced by the presence of drugs that act as antitumor agents. Topoisomerase inhibitors poison the cleavage intermediates and lead to the formation of an aborted or irreversible TOP2cc [39,47].

In order to repair the TOP2cc captured by the TOP2 poisons, the non-homologous end joining (NHEJ) or homologous recombination (HR) repair pathway is activated. To proceed with NHEJ repair, the trapped TOP2cc is first degraded by the proteasome, and then the remaining tyrosine-linked end can be released by tyrosyl-DNA phosphodiesterase 2 (TDP2) [59,60]. The TOP2cc (which is trapped by etoposide), is captured by TDP2, the latter hydrolyzes the phosphodiester bond between TOP2 and DNA through proteasome degradation [61]. The collaboration between TDP2 and proteasome is not well understood.

The study by Schellenberg et al. provided some novel findings [62]. First, zinc finger protein 451 (ZNF451) binds and reshapes TOP2cc, thereby opening the complex and allowing TDP2 to access and interact with the cleavage complex. Second, ZNF451 assists small ubiquitin like modifier 2 (SUMO2) protein in SUMOylation of trapped TOP2cc. The interaction of TDP2 with SUMO2 bound to TOP2cc further facilitates the interaction of TDP2 with the cleavage complex. This activity is instrumental in preventing the formation of the cell-deadly DSBR that is typical of TOP2 poisons. To proceed with HR repair, the MRE11-RAD50-NBS1 (MRN) complex cooperates with other repair proteins (e.g., BRCA1 and CtIP) and can directly sever a small segment of DNA ends containing covalently bound TOP2. In case of DNA damage is recognized, the MRN complex recruits and activates ataxia-telangiectasia mutant kinase (ATM) dimers [63]. The activated ATM dimer interacts with the checkpoint kinase, causing the cell cycle to stop at G2/M [64]. Subsequently, the nuclease activity of the MRN complex cleaves the DNA ends, prompting HR repair. Meanwhile, it was demonstrated that MRN, CtIP, and BRCA1 are required to remove TOP2-DNA adducts induced by etoposide treatment, and subsequent excision of TOP2-induced DSBR ends.

It is revealed that the interaction between CtIP and BRCA1 is necessary for the resistance of cells to etoposide during genomic DNA replication [65]. In addition, the MRN complex is also involved in the repair of NHEJ, and is species-specific in the repair of the hairpin structure, but the exact mechanism is still unclear [66]. The degradation of TOP2 through these two processes will leave free DNA ends, which undergo continuous excision and final repair [67,68].

Once TOP2 is released from the DNA-TOP2 cross-linking adduct, NHEJ or HR repair is initiated. NHEJ repairs DSBR directly rejoining DNA terminals, which may produce a perfect repair. If the sequence around the lesion is missing or presenting with other DSBR, deletion or translocation may occur [69]. Compared to NHEJ, HR is a more precise mechanism where a homologous sequence is used as a template to guide the repair process. The DSBR can be repaired by several homologous mediated pathways, including double-strand break repair (DSBR) and synthesis-dependent strand annealing (SDSA), both of the pathways are initiated by 3′-single-strand DNA [70]. The DSBR pathway produces a crossover or non-crossover product, while the SDSA pathway produces a non-crossover product. Crossover is essential for proper chromosome segregation during meiosis and interprets genetic variation. In fact, it does not rule out that a perfect repair can be produced if the template sequence is the same as the sequence with the break sequence. Rarely, if the template is a non-homologous sequence, it may produce a translocation as well [71]. NHEJ is usually involved in the development of lymphocytes, while HR is thought to be more involved in the catalytic recombination of meiosis [71,72], which suggests that the NHEJ pathway is mainly involved in DSBR repair induced by TOP2 poisons. In this regard, several studies have emphasized that NHEJ pathway plays a vital role in the TOP2 poison induced DSB (etoposide was specifically listed) [73,74] (Fig. 3).

Etoposide and chromosomal translocations

A distinct subset of t-AML is associated closely with TOP2 poisons. t-AML shows unique cytogenetic changes, the most common of which is the destruction of the MLL gene on chromosome 11q23 in the 8.3 kb breakpoint cluster region (BCR). The 7 to 13 exon of the MLL gene are located in the BCR, where most of the chromosomal translocation breakpoints are located [15,75]. Several lines of evidence suggested that etoposide caused TOP2cc cleavage activities induced site-specific DNA cleavage in the MLL BRC [76-78]. Likewise, Martin et al. reported that DNA cleavage within the MLL BCR is a specific event in the initial phase of apoptosis that is part of higher-order chromatin fragmentation leading to chromosomal translocations in the MLL and AML1 genes. In addition, in vitro incubation of etoposide with TOP2 enhanced the DNA-TOP2 cleavage complex near the translocation site within the DNA sequence containing MLL and partner genes identified in susceptible leukemia [77,79,80]. The partner genes found in MLL-related translocations include AF9, AF4, ENL, AF10, AF6, AF17, EPB1S, GAS7, LOC100128568, CREBBP, and PTD [81].

Etoposide metabolites contribute to therapy-related leukemia: TOP2 poison

As described above, etoposide metabolites display activity against TOP2 enzymes, which is similar or greater than etoposide [82-85]. However, studies have shown that etoposide quinone or etoposide catechol has a similar inhibitory activity towards TOP2, when performed with a buffer containing a large amount of reducing agents [26,86,87]. Studies with minimal reducing agents have shown that etoposide quinone is several times more effective than the parent compound in inducing TOP2-mediated DSBR [83,84]. It was simultaneously demonstrated that the inhibition of TOP2 by etoposide quinone is not limited in the same manner as etoposide by blocking ligation as an interfacial poison [37,82–84]. Numerous studies imply that etoposide quinone is a redox-sensitive compound [84,88–90]. More significantly, etoposide quinone was reported to act as a covalent poison, a mechanism involving covalent addition, which may lead to multiple effects: inactivation of the enzyme may occur due to blocking of DNA before binding to the enzyme or stabilizing N-terminal clasp after DNA binding to the enzyme [82]. In addition, etoposide quinone is a covalent poison of TOP2α and TOP2β, and its reactivity to TOP2β is reported to be slightly higher than that of TOP2α [85]. Meanwhile, studies have shown that etoposide catechol is 2–3 folds more potent than etoposide and can induce high levels of double-stranded DNA cleavage under oxidative conditions [83]. Compared with etoposide, its metabolites are several times more potent on TOP2, and they induce higher levels of DSBR under oxidative reaction conditions.

Benzene is a carcinogen that can induce AML in humans and various tumors in animals. According to several reports, benzene metabolites, including benzoquinone (BQ), are responsible for the genotoxic and leukemogenic effects of benzene . In particular, benzoquinone can directly modify DNA, causing genotoxicity and mutagenesis [91,92]. BQ is a TOP2 poison as well, similar to etoposide quinone, which blocks DNA linkage and produces TOP2-DNA adducts [47,93]. In addition, exposure to benzene metabolites in mouse fetal liver is known to increase reactive oxygen species (ROS) [94,95]. ROS has been shown to contribute to the modification of topoisomerases [95,96]. The c-Myb transcription factor has also been shown to be abnormally activated in hematopoietic cells exposed to BQ [98]. In addition, NAD (P)H quinone dehydrogenase 1 (NQO1) is an enzyme that metabolizes BQ into hydroquinone, which is a less reactive compound. C609T mutation encoding an inactive form of NQO1 has been suggested to increase the
risk of leukemia with chromosome 11q23 translocations [99,100] (particularly prominent in acute non-lymphoblastic leukemia, where the MLL gene is fused to the AF-4 gene on chromosome 4) [101].

Etoposide metabolites contribute to therapy-related leukemia: non-TOP2 factors

Exposure to benzene is correlated with the deregulation of the expression of specific genes associated with leukemia and DNA methylation changes [102,103]. Chromatin components, in particular histones may also react with etoposide (or its metabolite) as shown for histone H1 [104]. The benzene metabolite 1,2,4-benzenetriol can influence DNA methylation and histone acetylation in K562 cells [105]. Recently, etoposide quinone has been shown to inhibit certain enzymes, such as the histone acetyltransferase CREBBP which is known to be involved in the leukemogenesis [106]. CREBBP (also known as CBP or KAT6A) regulates both normal and malignant hematopoiesis [107], and the majority of Crebbp-/- or Crebbp-/- mice develop malignant hematopoietic pathologies indicating that CREBBP acts as a tumor suppressor and plays a crucial role in hematopoiesis [108–110]. We recently showed that etoposide quinone can inhibit CREBBP acetyltransferase activity by reacting with cysteine within zinc fingers domains that are key for CREBBP activity [106].

TCPTP (also known as PTPN2) is a member of the protein tyrosine phosphatase (PTP) family, which negatively regulates JAK/STAT (Janus kinase/signaling transducer and activator of transcription) signaling, through dephosphorylation of different tyrosine phosphorylated JAK/STAT proteins, for instance, STAT1 or JAK1. TCPTP plays an essential role in normal and malignant hematopoiesis by regulating the JAK/STAT pathway [111,112]. Deletions or inactivating mutations of TCPTP were identified in non-Hodgkin’s lymphoma and T-cell leukemia and associated with elevated STAT signaling and changes in gene expression.

Fig. 3. TOP2 poison induces translocation mediated by the DSB repair pathway. TOP2 poison-mediated DSB is likely to be repaired by two pathways, NHEJ or HR repair. NHEJ can repair precisely when the DSB at a low concentration level. If DSB accumulates to a high concentration level, it may cause sequence deletion and translocation. NHEJ mainly contains three types of repair products, faithful repair, sequence loss, and translocation. HR is mediated by a homologous sequence. According to the model sequence, it may produce a different product. In most cases, HR repair uses homologous sequences as templates. This situation generates mainly a crossover or a non-crossover product through two pathways, including SDSA and DSBR, which play an instrumental role in meiosis. In a few cases, precision repair occurs when HR repair uses the same sequence as a template, and translocations may happen when HR uses a non-homologous sequence as a template. The blue and the red chromosomes are homologous, while the orange indicates a non-homologous chromosome. The blue arrowheads are located at one of the Holliday junctions (HJ), and the red arrowheads are located at the other junctions.
The loss of TCPTP phosphatase activity may lead to excessive activation of the JAK/STAT pathway, which further changes the development of HSC. Etoposide and benzene metabolites, including etoposide quinone and benzoquinone, can irreversibly inhibit TCPTP activity, suggesting another new mechanisms for etoposide and benzene to induction of leukemia [115,116]. (Fig. 4)

Conclusion and further challenge

In summary, etoposide and its metabolite etoposide quinone can inhibit TOP2 enzymes, and the accumulation of DSB is the main cause of secondary leukemia. There is still a need to obtain more precise evidence to prove that CREBBP and TCPTP are involved treatment-related leukemia. However, it has been established that etoposide quinone could contribute more importantly to the development of leukemia than etoposide through TOP2 inhibition. It is likely that other mechanisms such as CREBBP and TCPTP inhibition, may also contribute to etoposide treatment-related leukemia.

Similarly, a large amount of evidence has been established that metabolites of benzene are involved in leukemogenesis through other mechanisms than TOP2 inhibition. For instance, chronic occupational exposure to benzene leads to an increase in the expression of genes involved in apoptosis in blood cells [117,118]. Benzene metabolite triggers apoptosis of hematopoietic progenitor cells in a dose-dependent manner in bone marrow [119]. Meanwhile, benzene affects cell apoptosis by inhibiting caspase-3 [120,121], which may promote the survival of cancer cells. It is also well known that inappropriate activation or expression of c-Myb is involved in the process of leukemia. The c-Myb transcription factor has also been shown to be abnormally activated in K562 and mouse models of hematopoietic cells exposed to 1,4-benzoquinone [122,123]. Moreover, the duration of exposure to benzene is associated with the dysregulation of specific gene expression associated with leukemia and changes in DNA methylation [102,103].

Based on the existence of homologous histone acetyltransferases or protein tyrosine phosphatases, it is likely that etoposide quinone may affect homologs of CREBBP and TCPTP. For example, p300, the homolog of CREBBP, is structurally and functionally similar to CREBBP, notably, on the acetyltransferase catalytic core, including RING and PHD domains. Etoposide quinone covalently binds to CREBBP in the RING and PHD domains [106]. Indeed, p300 has similar RING and PHD domains, it is thus likely that etoposide quinone may also inhibit the acetyltransferase activity of p300. Other histone acetyltransferases such as TIP60, MOZ, and MOF, contain structural zinc fingers and may also be targets of etoposide quinone. Acetyltransferases, particularly CREBBP/p300, are involved in HR repair by acetylating Recombinant DNA repair protein 52 (RAD52) [124,125]. TIP60 plays a direct role in

Fig. 4. The underlying mechanism of etoposide induced therapy-related leukemia. TOP2 is an essential as well as a genotoxic enzyme. Typically, the TOP2, DNA, and TOP2-DNA complex are maintained precisely to perform the relevant cellular functions. If the TOP2 enzyme activity is insufficient, the TOP2-DNA complex is lacking, resulting in the inability to release the torsional stress, leading to cell division failure and eventual death due to the failure to unwind the daughter chromosomes. In contrast, TOP2 poisons inactivate the enzymes via interfacial or covalent interaction, resulting in irreversible damage to the TOP2-DNA complex and ultimately creating DSB. Accumulation of DSB inhibits fundamental DNA processes and initiates recombination/repair pathways that produce chromosomal translocations and mutations. If the DSB overwhelms the cells, they will trigger apoptosis, which is the primary anticancer mechanism of etoposide. If the concentration of TOP2 mediated DSB is limited to induce apoptosis, mutations or chromosomal aberrations could occur, and it is possible to develop cancers. Etoposide caused therapy-related leukemia is a chromosomal rearrangement involving the MLL gene on chromosome 11q23. Meantime, etoposide quinone inhibits CREBBP or TCPTP, which has been reported to be associated with leukemogenesis.
activating ATM (an essential DBS repair protein kinase) by acetylating it in response to DNA damage [126,127]. Therefore, the loss of acetyltransferase activity may directly affect DBS repair.

The catalytic cystine 216 is an essential residue of TCPTP and participates to the enzymatic reaction. Cysteine 216 is a evolutionary conserved residue in the PTP family [128]. Therefore, etoposide quinone may inhibit other PTP family members, possibly leading to a broad loss of PTP activity in cells. This may affect JAK/STAT signal transduction which plays a key role in HSC differentiation and proliferation. In parallel, a study reported that deficiency in protein tyrosine phosphatase 1B (PTP1B) is involved in the development of acute leukemia [129]. In addition, SHP2 (Src homology region 2 domain containing phosphatase 2, also known as PTPN11) is related to hematopoiesis and leukemia [130].

As suggested for benzene-induced leukemia, etoposide-induced secondary leukemia are likely to rely on different mechanisms of actions affecting several and distinct biological processes. More studies are needed to understand the pathogenesis of etoposide therapy-related leukemia. This may facilitate the determination of the optimal therapeutic administration methods, including the timing and dosage and thereby advances the prevention and treatment of etoposide therapy-related leukemia.

Declaration of Competing Interest

All authors declare no conflicts of interest.

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Author contributions

WZ, and PG designed the review. JMD, CC and FRL provided expertise. All authors contributed intellectually and to the writing of the manuscript.

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