ORIGINAL RESEARCH

Characterization of Stable and Reactive Metabolites of the Anticancer Drug, Ensartinib, in Human Liver Microsomes Using LC-MS/MS: An in silico and Practical Bioactivation Approach

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Department of Pharmaceutical Chemistry, College of Pharmacy, King Saud University, Riyadh 11451, Kingdom of Saudi Arabia **Background:** Ensartinib (ESB) is a novel anaplastic lymphoma kinase inhibitor (ALK) with additional activity against Abelson murine leukemia (ABL), met proto-oncogene (MET), receptor tyrosine kinase (AXL), and v-ros UR2 sarcoma virus oncogene homolog 1 (ROS1) and is considered a safer alternative for other ALK inhibitors. ESB chemical structure contains a dichloro-fluorophenyl ring and cyclic tertiary amine rings (piperazine) that can be bioactivated generating reactive intermediates.

Methods: In vitro metabolic study of ESB with human liver microsomes (HLMs) was performed and the hypothesis of generating reactive intermediates during metabolism was tested utilizing trapping agents to capture and stabilize reactive intermediates to facilitate their LC-MS/MS detection. Reduced glutathione (GSH) and potassium cyanide (KCN) were utilized as trapping agents for quinone methide and iminium intermediates, respectively.

Results: Four in vitro ESB phase I metabolites were characterized. Three reactive intermediates including one epoxide and one iminium intermediates were characterized. ESB bioactivation is proposed to occur through unexpected metabolic pathways. The piperazine ring was bioactivated through iminium ions intermediates generation, while the dichlorophenyl group was bioactivated through a special mechanism that was revealed by LC-MS /MS.

Conclusion: These findings lay the foundations for additional work on ESB toxicity. Substituents to the bioactive centers (piperazine ring), either for blocking or isosteric replacement, would likely block or interrupt hydroxylation reaction that will end the bioactivation sequence.

Keywords: in vitro phase-I metabolites, iminium intermediates, epoxide intermediates, human liver microsomes

Introduction

Cancer is the second leading cause of death worldwide and is accountable for about 9.6 million casualties in 2018, with almost one-sixth of global deaths.¹ Earlier studies have shown that molecular-targeted therapeutic approaches were successful in treating cancer cases by targeting genes, oncogenes, and tumor suppressors involved in cancer progression.² Lung cancer has the highest mortality rate globally among all cancer types. In 2012, 1.59 million deaths were attributed to lung cancer, which is 20% of all cancer deaths.³ Non-small-cell lung cancers (NSCLCs) account

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First-generation ALK inhibitors (eg, crizotinib) became less active due to NSCLC resistance mutations; however, new-generation ALK inhibitors (eg, brigatinib and ensartinib (ESB)) were designed to treat these mutations.^{12,13} ESB is a novel ALK inhibitor with additional activity against ABL, MET, ROS1, and Axl. It provided promising preliminary data in terms of safety and clinical activity.^{14,15} In phase I/II clinical trial, ESB was well tolerated and active in patients with ALK-positive NSCLC¹⁶ and showed intracranial responses in patients with ALK-positive NSCLC who developed CNS metastases.¹⁷

Drug detoxification occurs through metabolic pathways that improve the hydrophilicity of xenobiotics and endogenous compounds facilitating their elimination outside the human body. Additionally, compared with the precursor molecules, the produced metabolites demonstrated lower toxicity profiles. Nonetheless, in special cases, reactive metabolites (unstable intermediates) were produced exhibiting greater toxicity.¹⁸⁻²⁰ These unstable intermediates are considered electron-deficient species, so covalent bonds can alter DNA and proteins, which is the preliminary step in organ toxicity.^{21,22} Characterization of the generation of reactive intermediates is crucial to understanding drug-induced toxicity. Reactive intermediates are often generated through metabolic pathways in phase I metabolism and cannot be identified by direct methodology as they are unstable species. Thus, nucleophile trapping can be utilized to seize the generated reactive species forming stable adducts that could be characterized using liquid chromatography-tandem mass spectrometry (LC-MS/MS).²³⁻²⁶

The IUPAC designation of ESB is 6-amino-5-[(1R)-1-(2,6-dichloro-3-fluorophenyl) ethoxy]-N-(4-{[(3R, 5S)-3,5-dimethyl-1-piperazinyl] carbonyl} phenyl)-3-pyridazinecarboxamide (Figure 1). ESB chemical structure contains a dichloro-fluorophenyl ring and cyclic tertiary amine rings (piperazine). Human liver



Figure I Chemical structure of ensartinib.

microsomes (HLMs) can generate unstable iminium while metabolizing piperazine that can be trapped using potassium cyanide (KCN).²⁷⁻³⁰ The dichlorofluorophenyl ring can be bioactivated to an unstable epoxide species by oxidative defluorination mechanism generating quinone methide intermediate that can be captured by reduced glutathione (GSH).³¹ The formed stable adducts, which were formed by nucleophilic-electrophilic reactions, can be separated by highperformance liquid chromatography (HPLC) and characterized by mass spectrometry (MS) analyzers.^{23,24,27,32,33} These reactive electrophiles may contribute to the ESB reported side-effects.

In patients receiving ESB treatment, common toxicities were as follows: vomiting (26%), fatigue (22%), nausea (36%), rash (56%), and pruritus (28%), with about 23% of the patients exhibiting a grade 3–4 treatment-related toxicity (including pruritus and rash).^{17,34} Here, LC-MS/MS was employed to characterize in vitro phase I metabolites of ESB generated in the incubation of HLMs. The generation of reactive electrophiles was also tested using capturing agents. Notably, ESB bioactivation was observed to take place via unexpected pathways.

Piperazine ring was bioactivated forming unstable iminium ion species. The dichloro-3-fluorophenyl ring was bioactivated by an unexpected metabolic reaction verified by LC-MS/MS. ESB generated three reactive metabolites through three ways of bioactivation that were detected via in silico testing and in vitro phase I metabolic practical experiments. This could be a new strategy for reducing the side-effects of the newly designed drugs without affecting and even improving its pharmacological activity. Targeted modifications for the characterized structural alerts and bioactive centers may improve the safety of ESB while maintaining its efficacy, which was confirmed by the DEREK module for StarDrop software. Four in vitro phase I metabolites, one GSH, and two cyano adducts of ESB were characterized. The proposed reaction involved oxidation, hydroxylation, and cyanide or GSH addition. In silico testing of the toxicity of ESB-related metabolites was performed using DEREK software. To the best of our knowledge, this is the first experiment of in vitro phase I metabolites of ESB, including structural characterization of novel ESB reactive intermediates.

Chemicals and Methods

Chemicals and Instruments

HLMs (product number: M0567), collected from various male human livers, were obtained from Sigma-Aldrich (St. Louis, Missouri, USA) and stored at -70°C until use. Organic solvents and reference powders were of HPLC grade and analytical grade (AR), respectively. HPLC grade water was prepared in-house by a Milli-Q Plus filtration instrument (Millipore, USA). ESB (98.51%) and NADPH (99.99%) were obtained from MedChemExpress (USA). Acetonitrile, formic acid, ammonium formate, GSH, and KCN were acquired from Sigma-Aldrich (USA). LC-MS/MS analysis was

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performed using Agilent 1200 HPLC connected to Agilent 6410 triple quadrupole. Electrospray ionization (ESI) was used as an interface. Incubation of ESB with HLMs was performed in the SW22 Shaking Water Bath (JULABO, Seelbach, Germany) adjusted at 37°C.

Chromatography Conditions

Both liquid chromatographic and mass optimized conditions for resolution and identification of the ESB-HLM incubation mixture are provided in Table 1.

In silico Prediction of ESB Metabolites Using WhichP450™ Software

In this study, we aim to recognize the susceptibility of metabolism key sites, demonstrated by decreasing the site vulnerability, which is revealed by the composite site lability (CSL) scores, and examine the proposed regioselectivity of major isoforms responsible for metabolism. The outcomes from the WhichP450[™] module of StarDrop software, shown in a pie chart, were used as an indication of the most likely CYP450 isoform that plays an important role in ESB metabolism.

| Agilent RRLC | | | Triple Quadrupole MS | |
|---|---|-----|----------------------|-----------------------------------|
| Model | Agilent 1200 | | Model | Agilent 6410 |
| Gradient mobile phase | A: Aqueous (5 mM NH ₄ COOH in H ₂ O at pH: 3.8) | | ESI | Positive ESI mode |
| | B: Organic (Acetonitrile) | | | Drying nitrogen gas |
| | 0.3 mL/min | | | 11 L/min flow rate 55 psi |
| | 50 min | | | |
| C ₁₈ column (Agilent Eclipse Plus) | s) 250 mm length 4.6 mm ID 5 μm particle size 21 ± 2°C | | | 350°C |
| | | | | 4000 V capillary voltage: |
| | | | Analyzer modes | MS full scan and PI positive mode |
| | | | Collision gas | Nitrogen (High-purity) |
| Gradient mobile phase | Time in min | % B | Analytes | ESB and its metabolites |
| | 0 | 5 | | |
| | 5 | 5 | Analyzer | 25 eV Collision energy |
| | 30 | 60 | | |
| | 45 | 90 | | |
| | 50 | 5 | 1 | 150 eV Fragmentor voltage |

 Table I LC-MS/MS Optimized Chromatographic Parameters



Figure 2 Suggested metabolic labile sites for ensartinib (ESB) by CYPP450 module of StarDrop software.



Figure 3 Outcomes of in silico reactivity tool expectation by the Xenosite including Cyanide and GSH (A). Structural alerts of ESB as proposed by DEREK toxicity predictor software (B).

| Code | Skin Sensitization | Carcinogenicity | Nephrotoxicity | Chromosome Damage, | |
|------|----------------------|---------------------------------------|------------------------|--|--|
| | Activated benzene | Polyhalogenated aromatic or Alkylaryl | Halogenated benzene | Genotoxicity, Teratogenicity and Mutagenicity | |
| ESB | Plausible | Plausible | Equivocal | NA | |
| MI | Plausible | Plausible | Equivocal | NA | |
| M2 | Plausible | Plausible | Equivocal | NA | |
| M3 | Plausible | Plausible | Equivocal | NA | |
| M4 | Plausible | Plausible | Equivocal | NA | |

 Table 2 Qualitative Toxicity Prediction (DEREK) of the ESB and Its Metabolites

In silico Prediction of ESB Reactive Metabolites Using XenoSite Reactivity Model and DEREK Software

XenoSite reactivity model in silico studies, freely available online at <u>http://swami.wustl.edu/xenosite</u>, were conducted to detect liable sites that can produce reactive intermediates forming stable adducts with GSH and cyanide.³⁵ This prognostic model was built on neural networks of more than 680 molecules. It has a short processing time.^{36,37} The SMILES format of the ESB chemical structure was uploaded to the XenoSite reactivity model online website. DEREK software was employed to check for structural alerts, verify our proposed bioactivation pathway, and propose structural modification at the bioactive sites that may stop the toxicity sequence.

HLM Incubation

ESB was solubilized in dimethyl sulfoxide (DMSO). To avoid nonspecific protein binding, 1 mg/mL of HLMs was used. The characterization for ESB-related metabolites was performed by incubation of ESB (30μ M) with HLMs (1.0 mg/mL) in 1 mL of 50 mM phosphate buffer (pH, 7.4) plus 3.3 mM MgCl₂ in a thermostatic shaking water bath (37° C) for 2 h. Incubation was initiated by adding NADPH to a final concentration of 1.0 mM after 5 min of preincubation and terminated by adding 2 mL of ice-cold ACN as a precipitating agent and extraction solvent. Excess precipitated protein was discarded by centrifugation at 9000 g for 15 min at 4°C. The purified supernatants were transferred to clean tubes; then,



Figure 4 Fragment ion (m/z 561) chromatogram displaying the ESB peak at 24.0 min (A). Fragment ions of ESB (B).



Scheme I ESB fragment ions.

evaporation to dryness was carried out under the influence of a nitrogen stream. The evaporated samples were reconstituted in the mobile phase; then, they were transferred to HPLC vials for LC-MS/MS analysis (Agilent 6410 QqQ).^{38–41} The ESB-HLM incubation was repeated using the same procedure by adding GSH or KCN for capturing reactive metabolites. All metabolic procedures were conducted three times using controls (without NADPH or HLMs) to confirm that all identified stable and reactive metabolites were metabolically generated.

Identification of ESB Reactive Metabolites

After a full mass scan (MS) over a specific m/z range, extraction of molecular ion peaks (MIP) from the total ion chromatograms (TIC) was utilized to find the hypothesized

metabolites. Chromatographic peaks (CP) of the predicted metabolites were exposed to fragmentation using high purity nitrogen under the collision-induced dissociation (CID) mode within the collision cell of the triple quadrupole, resulting in the generation of fragment ions required to reconstruct the chemical structures into the proposed chemical structure. Thus, the fragmentation pattern of ESB was used as a confirmatory and interpretation tool for the predicted chemical structures of the in vitro stable and reactive metabolites generated during ESB metabolism.

Results and Discussion In silico Prediction of ESB Metabolites

The ESB metabolic landscape exhibits the vulnerability of each chemical site towards CYP3A4 metabolism in

| Ensartinib | Full MS Scan | Qualitative Fragment Ions | Elution Time (Min) | Proposed Metabolic Pathways |
|---------------------|-----------------|------------------------------|-----------------------|--|
| | 561 | 447, 257, 138 | 24.0 | Main Drug |
| Phase-I metabolites | | | | |
| MI | 575 | 447, 368, 257 | 20.9 | Oxidation at piperazine ring |
| M2 | 575 | 503, 447, 257, 203, 85, | 23.3 | Oxidation at methyl group attached to piperazine |
| | | 57 | | ring |
| M3 | 577 | 463, 371, 327, 257, 120 | 18.9 | Hydroxylation at benzoyl ring |
| M4 | 577 | 447, 257, 147 | 21.5 | Hydroxylation at piperazine ring |
| Trapped reactive | | | | |
| intermediates | | | | |
| ESB602 | 602 | 557, 447, 257, 138 | 26.5 | Hydroxylation and cyano addition at piperazine |
| | | | | ring |
| ESB866 | 866 | 848, 790, 496, 371, 257 | 16.8 | Defluorination, hydroxylation, oxidation and GSH |
| | | | | conjugation |

Table 3 In vitro Metabolites and Reactive Intermediates of Ensartinib



Figure 5 Fragment ion (m/z 575) chromatogram displaying the MI and M2 peaks at 20.9 min and 23.3 min, respectively (A). Fragment ions of MI (B). Fragment ions of M2 (C).

absolute terms, to guide the ESB metabolites prediction and the chemical structure optimization for enhancing metabolic stability. This reveals that dimethyl piperazine ring and dichlorophenyl group are labile which is consistent with experimental laboratory work. The CSL is shown in the metabolic landscape at the top-right corner and the P450 column of the dataset (Figure 2). CYP3A4 was predicted to play an important role in ESB metabolism. CSL value is 0.98 indicating high liability for ESB to metabolism. The metabolic liability of ESB is mainly



Scheme 2 MI and M2 fragment ions.

due to N7, C1, C2, C6, and C8 of substituted methyl piperazine ring.

In silico ESB Bioactivity and Toxicity Prediction

The vulnerable electrophilic atomic sites of ESB were proposed for cyanide and GSH attack and conjugation. The outcomes were represented by a color scale bar that has a color gradient as follows: the red color indicates the maximum possibility of reactive intermediate generation at the site, whereas the white color indicates no possibility of reactive intermediate formation (Figure 3A). The cyanide model indicates that the piperazine ring could be metabolized generating reactive iminium intermediates that could be attacked by cyanide ion forming cyano adduct. Furthermore, dichlorophenyl and pyridazine rings were predicted for GSH conjugation. The faint blue color means that forming reactive intermediates are less probable, which was confirmed by in vitro experiments. The outcomes of the predictions are shown in Figure 3A.

As proposed by DEREK software (in silico toxicity evaluation program), ESB and related metabolites were expected to exhibit skin sensitization, nephrotoxicity, and carcinogenicity due to activated benzene, halogenated benzene, and polyhalogenated aromatic, respectively, which were toxicity alerts by the software (Figure 3B). On the other hand, M1, M2, and M3 were expected to exhibit carcinogenicity due to alkyl aryl moiety. Table 2 lists the in vitro ESB stable metabolites with DEREK results for the predicted toxicity profile (Figure 3B).



Figure 6 Fragment ion (m/z 575) chromatogram displaying the M3 and M4 peaks at 18.9 min and 21.5 min, respectively (A). Fragment ions of M3 (B). Fragment ions of M4 (C).



Scheme 3 M3 fragment ions.



Scheme 4 M4 fragment ions.



Figure 7 Fragment ion (m/z 602) chromatogram displaying the ESB602 peak at 26.5 min (A). Fragment ions of ESB602 (B).



Scheme 5 ESB602b fragment ions.

ESB Dissociation Pattern

The ESB CP appeared at 24.0 min in the TIC (Figure 4A). CID of the ESB at m/z 561 resulted in the generation of three qualitative fragment ions at m/z 447, 257, and 138, showing weak chemical bonds in the ESB structure (Figure 4B, Scheme 1). The fragment ion at m/z 447 reveals the loss of the dimethyl piperazine ring, whereas the other two fragment ions at m/z 138 and m/z 257 have qualitative significance in the structural rebuilding of the related metabolites.

Identification of ESB Stable and Reactive Metabolites

Four ESB metabolites were formed by two phase I metabolic pathways (oxidation and hydroxylation). One GSH adduct and two cyano adducts were characterized (Table 3). One GSH conjugate and two cyano adducts were identified after HLM incubation of ESB in the presence of trapping agents (GSH or KCN). Compared to CID of ESB, neutral loss of HCN (27 m/z) or fragment ion with 25 m/z more was indicative of a cyano group addition.



Figure 8 Fragment ion (m/z 866) chromatogram displaying the ESB866 chromatographic peak at 16.8 min (A). Fragment ions of ESB866 (B).

Scheme 6 ESB866 fragment ions.

MI and M2 Fragmentation Patterns

The ESB575 CPs appeared at two elution times (20.9 min and 23.3 min) in the TIC (Figure 5A). The CID of M1 and M2 ions at m/z 575 resulted in the generation of different fragment ions for the two peaks, whereas the CID of M1 and M2 ions at m/z 447 and 257 led to the generation of two qualitative fragment ions (Figure 5B and C). Compared to the CID of ESB, the fragment ions at m/z 447 and 257 were indicative of the oxidation metabolic pathway occurring at the piperazine moiety without knowing the exact place of hydroxylation, so a Markush structure was used to indicate the ambiguous structural assignment (Scheme 2). Oxidation may occur at α -carbon atoms to the tertiary nitrogen of piperazine moiety or the methyl group attached to the piperazine ring.

M3 and M4 Fragmentation Patterns

M3 and M4 CPs appeared at two elution times (18.9 min and 21.5 min), respectively, in the TIC (Figure 6A). The CID of M3 and M4 ions at m/z 577 resulted in the

Scheme 7 Supposed metabolic sequence for reactive iminium intermediate formation at piperazine rings in ESB metabolism and a potential capturing strategy.

Scheme 8 Supposed pathways of GSH conjugation through oxidative defluorination mechanism generating quinone methide intermediate.

generation of various fragment ions for the two peaks. The CID of M3 ion generated five fragment ions at m/z 463, 371, 327, 257, and 120 (Figure 6B). Compared to the CID of ESB, the fragment ions at m/z 463 (16 m/z more) were indicative of the hydroxylation metabolic pathway occurring at phenyl ring confirmed by the four fragment ions at m/z 371, 327, 257, and 120 (Scheme 3). The CID of M4 ion resulted in the generation of three fragment ions at m/z 447, 257, and 147. The exact place of hydroxylation at the phenyl ring is not known; thus, a Markush structure was used to detect the ambiguous structural assignment (Figure 6C). Compared to CID of ESB, the fragment ions at m/z 447 and 257 were indicative of the hydroxylation; without knowing the exact place of hydroxylation;

therefore, a Markush structure was used to indicate the ambiguous structural assignment (Scheme 4). Hydroxylation may occur at α -carbon atoms to the tertiary nitrogen of piperazine moiety or the methyl group attached to the piperazine ring (Scheme 4).

ESB602a and ESB602b Fragmentation Patterns

The ESB602 CP appeared at the elution time (26.5 min) in the TIC (Figure 7A). ESB602 molecular ion fragmentation resulted in the formation of three qualitative fragment ions at m/z 557, 447, and 257 (Figure 7B). Compared to CID of ESB, the fragment ions at m/z 447 and 257 proposed that all metabolic pathways occurred at the dimethyl piperazine moiety, keeping the remaining structure unchanged. DI at 557

Figure 9 Chemical structure of ensartinib displaying various metabolic phase I pathways and the proposed bioactive sites.

represented the neutral loss of one molecule of water and one molecule of HCN, revealing hydroxylation and cyano addition at the piperazine ring. The hydroxylation metabolic reaction and cyano addition occurred at the two α -carbon atoms to the tertiary nitrogen at the piperazine ring (Scheme 5).

ESB866 Fragmentation Pattern

The ESB866 CP appeared at 16.8 min in the TIC chromatogram (Figure 8A). CID of ESB866 ion at m/z 866 resulted in the generation of five fragment ions at m/z 848, 790 496, 371, and 257 (Figure 8B). Compared to the CID of ESB, the fragment ions at m/z 371 and 257 indicated the metabolic bioactivation and then conjugation with GSH at the dichlorophenyl ring. We proposed that epoxide was formed by the direct oxidation of the dichlorophenyl ring generating reactive intermediate (quinone methide) attacked by the GSHforming adduct (Scheme 6). This mechanism of metabolic bioactivation has been discussed for other compounds (eg, X-376 and diclofenac).^{31,42,43}

ESB Bioactivation Sequence

ESB602 cyanide adduct characterization revealed the formation of unstable iminium metabolites in the in vitro metabolism of ESB and HLMs. The metabolic bioactivation process was initiated by hydroxylation metabolic reaction at the bioactivated piperazine ring in ESB followed by water loss (dehydration), which generated unstable metabolites. These unstable metabolites (reactive intermediates) could be stabilized by cyanide nucleophile forming cyano adducts (ESB602) (Scheme 7). The proposed pathway of the iminium intermediate formation in ESB in vitro metabolism has been previously discussed in the literature.^{44–47}

ESB866 GSH conjugate revealed the formation of reactive intermediate through oxidative defluorination process in the ESB in vitro metabolism generating quinone methide intermediate that was captured by GSH (Scheme 8).³¹ We proposed that epoxide was formed by the direct oxidation of the dichlorophenyl ring generating reactive intermediate that was attacked by GSH-forming adducts. The formation of GSH conjugate was confirmed using negative precursor ion mode (272) and positive neutral loss (129) mode of triple quadrupole mass analyzer.

Conclusions

Two metabolic reactions (oxidation and hydroxylation) generated four stable phase I ESB metabolites in the in vitro metabolism of ESB. The bioactivation sequence

resulted in the generation of one GSH conjugate and one cyano adduct (Figure 9). The piperazine ring was metabolically bioactivated due to the generation of an unstable reactive intermediate that may be responsible for ESB sideeffects. The dichlorophenyl ring was bioactivated by oxidative defluorination pathway generating quinone methide intermediate, which was verified using LC-MS/MS. These findings provide the basis for further studies on ESB toxicity. Substituents to the bioactive centers (piperazine ring) for either blocking or isosteric replacement would likely block or interrupt the hydroxylation reaction that will end the bioactivation sequence. Future drug discovery studies can be conducted following this concept, which allows designing novel drugs with more safety profile without influencing their pharmacological effects.

Abbreviations

ESB, ensartinib; ESI, electrospray ionization; HLMs, human liver microsomes; TKIs, tyrosine kinase inhibitors; LC-MS/MS, liquid chromatography tandem mass spectrometry; GSH, reduced glutathione; KCN, potassium cyanide; NSCLCs, Non-small cell lung cancers; ALK, Anaplastic lymphoma kinase; CSL, composite site lability; HPLC, high-performance liquid chromatography; ABL, Abelson murine leukemia; MET, met proto-oncogene; ROS1, v-ros UR2 sarcoma virus oncogene homolog 1; Axl, receptor tyrosine kinase.

Ethics

The study's design (in vitro experiments using commercially available human liver microsomes) exempts it from the need of the Ethics Committees approval.

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Author Contributions

All authors contributed to data analysis, drafting or revising the article, have agreed on the journal to which the article will be submitted, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure

The authors report no conflicts of interest for this work.

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