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Overcoming acquired resistance of EGFR-mutant NSCLC cells to the third generation EGFR inhibitor, osimertinib, with the natural product honokiol

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1. Introduction

Lung cancer is the leading cause of cancer death worldwide and thus remains a serious health issue and economic burden (Torre et al., 2016). In recent years, targeted therapies have transformed the treatment of lung cancer patients with a specific mutation. Among them, epidermal growth factor receptor-tyrosine kinase inhibitors (EGFR-TKIs) benefit lung cancer patients with activating EGFR mutations (Tarapore and Lerose, 2015). This group of drugs has been developed rapidly from the initial 1st generation (e.g., gefitinib

Abbreviations
CHX, cycloheximide; EGFR, epidermal growth factor receptor; EGFR-TKIs, EGFR-tyrosine kinase inhibitors; HNK, honokiol; KO, knockout; NSCLC, nonsmall cell lung cancer; Osim, Osimertinib.
and erlotinib) to 2nd generation (e.g., osimertinib (Osim); also named AZD9291 or TAGRISSO™) EGFR-TKIs (Russo et al., 2017). However, the emergence of acquired resistance in patients occurs more rapidly than new-generation agents can be developed, and eventual treatment failure is inevitable (Russo et al., 2017).

The appearance of the T790M resistance mutation is the major mechanism accounting for ~60% of relapse cases from 1st generation EGFR-TKI treatment (Juchum et al., 2015; Remon et al., 2014; Tartarone and Lerose, 2015). To overcome this mutation, Osim and other 3-rd generation EGFR-TKIs were developed, which selectively and irreversibly inhibit EGFR carrying the common ‘sensitive’ mutations, 19del and L858R, and the resistant T790M mutation without affecting wild-type EGFR. Osim is an FDA-approved drug for the treatment of NSCLC harboring activating EGFR mutations (first-line) or that has become resistant to 1-st generation EGFR-TKIs through the T790M mutation (second-line). Without exception, patients also develop resistance to Osim, preventing long-term clinic remission (Govindan, 2015; Ramalingam et al., 2017). Hence, the development of effective strategies to overcome resistance to Osim and other 3-rd generation EGFR-TKIs is very important in the clinic setting.

Honokiol (HNK) is a natural product purified from the Magnolia tree and has been used in Chinese, Korean, and Japanese traditional medicine (Banik et al., 2019; Lee et al., 2011). Many preclinical studies have demonstrated the potential antitumor activity of HNK against different types of cancer such as breast cancer, neuroblastoma, bladder cancer, pancreatic cancer, and lung cancer (Averett et al., 2016; Hsiao et al., 2019; Lin et al., 2019; Pan et al., 2017; Sengupta et al., 2017). With the goal of developing efficacious combinatorial regimens to overcome Osim resistance, we have identified HNK as a potential agent that may sensitize Osim-resistant NSCLC cells to Osim. Thus, this study focused on demonstrating the activity of HNK and its derivatives, when combined with Osim, against the growth of different Osim-resistant cell lines in vitro and in vivo and understanding the underlying mechanisms.

2. Materials and methods

2.1. Reagents

The source and preparation of Osim and cycloheximide (CHX) were the same as described previously (Shi et al., 2017). HNK and its derivatives diethylstilbestrol (DEtS) and carbazole p-isomer (CAz-p) were described previously (Raja et al., 2008). Mcl-1, PARP, p-AKT (S473), AKT, p-ERK1/2 (T202/Y204), and ERK1/2 antibodies were purchased from Cell Signaling Technology, Inc (Beverly, MA, USA). Bim antibody was purchased from EMD Millipore (Burlington, MA, USA). GAPDH antibody was purchased from Proteintech, Inc (Rocky Hill, NJ, USA).

2.2. Cell lines and cell culture

The Osim -resistant cell lines PC-9/AR, PC-9/GR/AR, PC-9/3M, and HCC827/AR and culture conditions were the same as described previously (Shi et al., 2017). The PC-9 cell line with EGFR 19del and C797S double mutation (named PC-9/2M herein) used in a previous study (Niederst et al., 2015) was kindly provided by A. N. Hata (Harvard Medical School, Boston, MA, USA). H1975/OSIR was provided by Y. Yang (Mayo Clinic, Rochester, MN, USA). The features of these cell lines are summarized in Table 1.

Table 1. Osim-resistant EGFR-mutant NSCLC cell lines used in this study.

<table>
<thead>
<tr>
<th>Resistant cell line</th>
<th>EGFR mutations</th>
<th>MET alteration</th>
<th>Isogenic parental line</th>
</tr>
</thead>
<tbody>
<tr>
<td>First-line Osim resistance</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HCC827/AR</td>
<td>19del</td>
<td>Amplification</td>
<td>HCC827</td>
</tr>
<tr>
<td>PC-9/AR</td>
<td>19del</td>
<td>None</td>
<td>PC-9</td>
</tr>
<tr>
<td>PC-9/2M</td>
<td>19del, C797S (trans)</td>
<td>None</td>
<td>PC-9</td>
</tr>
<tr>
<td>H1975/OSIR</td>
<td>L858R, T790M</td>
<td>None</td>
<td>H1975</td>
</tr>
<tr>
<td>Second-line Osim resistance</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PC-9/GR/AR</td>
<td>19del, T790M</td>
<td>None</td>
<td>PC-9/GR</td>
</tr>
<tr>
<td>PC-9/3M</td>
<td>19del, T790M, C797S (cis)</td>
<td>None</td>
<td>PC-9</td>
</tr>
</tbody>
</table>
by infecting cells simultaneously with lentiviruses carrying CRISPR/Cas9 and SgBim for 48 h followed by puromycin selection for another 5 days as described previously (Qian et al. 2018). Single-cell clones were picked for screening for Bim knockout (KO) using western blotting.

2.3. Cell survival and apoptosis assays

Cells seeded in 96-well plates were treated the second day with the tested agents. After a given time period of treatment, viable cells were determined using sulforhodamine B (SRB) assay as described previously (Sun et al., 1997). The calculation of combination index for drug interaction (e.g., synergy) was performed using COMPUSYN software (Combo Syn, Inc., Paramus, NJ, USA). Cell apoptosis was detected with an annexin V/7-AAD apoptosis detection kit (BD Biosciences, San Jose, CA, USA) following the manufacturer’s manual. PARP was detected by western blotting as another indicator of apoptosis.

2.4. Western blot analysis

The procedures for the preparation of whole-cell protein lysates and western blot analysis were the same as we described previously (Shi et al., 2017; Shi et al., 2016).

2.5. Colony formation assay

The effects of the tested treatments on colony formation were conducted in 12-well plates as previously described (Shi et al., 2017; Shi et al., 2016).

2.6. Protein stability assay

Mcl-1 protein stability was determined using CHX chase assay as described previously (Koo et al., 2015; Shi et al., 2017).

2.7. Animal xenograft and treatments

Animal experiments, which were approved by the Institutional Animal Care and Use Committee (IACUC) of Emory University, were carried out as described previously (Shi et al., 2017). The treatments included vehicle control, Osim (10 mg·kg⁻¹·day⁻¹, 5 days-week⁻¹, og), HNK (50 mg·kg⁻¹·day⁻¹, ip), and their combination. We shared vehicle and Osim treatments with another study that examined the activity of LBH589 and Osim combination against Osim -resistant tumors (Li et al., 2019) in order to minimize the usage of animals. Tumor volumes were measured using caliper measurements and calculated with the formula \( V = \frac{\text{length} \times \text{width}^2}{2} \). At the end of the treatment, mice were weighed and euthanized with CO₂ asphyxia. The tumors were then collected and frozen in liquid nitrogen after weighting.

2.8. Statistical analysis

The statistical significance of differences between two groups was analyzed with two-sided unpaired Student’s t-tests when the variances were equal. Data were examined as suggested by the same software to verify that the assumptions for use of the t-tests held. Differences among multiple groups were analyzed with one-way ANOVA. \( P \) values < 0.05 were considered to be statistically significant.

3. Results

3.1. Combination of HNK and osimertinib synergistically decreases the survival of varied EGFR-mutant NSCLC cell lines with acquired resistance to Osim and inhibits colony formation and growth

We first determined the effect of Osim in the presence of HNK on the growth of different Osim -resistant EGFR-mutant NSCLC cell lines including PC-9/AR, PC-9-GR/AR, PC-9/2M (19del and C797S are trans located), PC-9/3M (19del, T790M, and C797S are cis located), HCC827/AR, and H1975/OSIR (Table 1). The combination of HNK and Osim was more active than either agent alone in decreasing the survival of these cell lines. The CIs were < 1, particularly when HNK at 10 µM was used, indicating synergistic effects on decreasing cell survival (Fig. 1A). The long-term colony formation assay also showed that the combination of HNK and Osim was more effective than either single agent alone in suppressing the formation and growth of colonies in several Osim -resistant cell lines (Fig. 1B). Clearly, the presence of HNK is able to resensitize these Osim-resistant cell lines to Osim.

3.2. Combination of HNK and osimertinib enhances induction of apoptosis in osimertinib-resistant cell lines

We next determined whether the combination effectively decreased the survival of these resistant cell lines through enhancing induction of apoptosis. As evaluated with annexin V/flow cytometry, the combination of...
Fig. 1. The combination of HNK and Osim synergistically decreases the survival and inhibits colony formation and growth of Osim-resistant EGFR-mutant NSCLC cell lines with augmented induction of apoptosis. (A) The indicated cell lines seeded in 96-well plates were treated the next day with the given concentrations of Osim alone, HNK alone, or their combinations. After 72 h, cell numbers were estimated using the SRB assay. The numbers inside the graphs are CIs for the given combinations. (B) The indicated cell lines were seeded in 12-well cell culture plates. On the second day, the cells were treated with fresh medium containing DMSO, 5 µM HNK alone, 200 nM Osim alone, and HNK plus Osim and the treatment was repeated every 3 days for a total of 12 days. (C, D) The indicated cell lines were exposed to DMSO, 10 µM HNK, 1 µM Osim, or HNK plus Osim for 72 h (C) or 48 h (D) and then harvested for the detection of apoptosis with annexin V/flow cytometry (C) and for the detection of PARP cleavage with western blotting (D). The data are means ± SDs of four replicates (A), triplicate (B), or duplicate (C) determinations. *P < 0.05, **P < 0.01, and ***P < 0.001 at least compared with other treatments. CF, cleaved form.

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HNK and Osim significantly induced apoptosis in comparison with each agent alone, which induced little or no apoptosis in PC-9/AR, PC-9/GR/AR, and PC-9/3M cells (Fig. 1C). Similarly, we detected much greater levels of cleaved PARP in all three tested cell lines exposed to the combination of HNK and Osim than in cells treated with either agent alone. The levels of intact or uncleaved PARP were also reduced much more in PC-9/GA/AR and PC-9/3M cells exposed to the combination than in cells treated with either single agent alone (Fig. 1D). Collectively, these results demonstrate that the combination of HNK and Osim enhances induction of apoptosis in Osim-resistant cell lines.

3.3. Combination of HNK and osimertinib enhances reduction in Mcl-1 levels through facilitating its degradation in osimertinib-resistant cells

To understand the molecular mechanism by which the HNK and AZD9291 combination augments induction of apoptosis, we examined its effects on the levels of Bim and Mcl-1, two key molecules involved in mediating apoptosis induced by Osim in sensitive EGFR-mutant NSCLC cells as we recently demonstrated (Shi et al., 2017). In PC-9/GR/AR cells, the combination of HNK and Osim was clearly more potent than each agent alone in increasing Bim levels and in decreasing Mcl-1 levels across the tested time period from 4 to 24 h (Fig. 2A). In two additional Osim-resistant cell lines, PC-9/AR and PC-9/3M, the combination also enhanced the reduction in Mcl-1 levels although the enhancement of Bim elevation was not observed after 12-h treatment (Fig. 2B). Collectively, it is clear that the combination of HNK and Osim universally enhances the reduction in Mcl-1 levels in Osim-resistant cell lines accompanied by enhanced Bim elevation in some cell lines.

We assessed the effects of the HNK and Osim combination on Akt and ERK phosphorylation in these resistant cell lines. The combination had no enhanced effects on p-Akt levels, but clearly augmented the decrease in p-ERK1/2 levels in comparison with single-agent treatment across the tested cell lines (Fig. 2C).

Since p-ERK suppression is associated with enhanced Mcl-1 degradation induced by Osim in sensitive EGFR-mutant NSCLC cells, as demonstrated recently (Shi et al. 2017), we then logically asked whether the HNK and Osim combination facilitates Mcl-1 degradation in Osim-resistant cells. By performing CHX assays, we observed that Mcl-1 was degraded more rapidly in both PC-9/GA/AR and PC-9/2M cell lines exposed to the HNK and AZD9291 combination than in those exposed to DMSO control (Fig. 2D), suggesting that the combination of HNK and Osim indeed enhances Mcl-1 degradation in Osim-resistant cells.

3.4. Enforced overexpression of Mcl-1 protects AZD9291-resistant cells from being killed by the combination of HNK and osimertinib

To determine the involvement of Mcl-1 reduction in mediating induction of apoptosis by the HNK and Osim combination, we introduced an Mcl-1 expression construct into PC-9/3M cells through lentiviral infection to generate the cell line PC-9/3M/Mcl-1 that expressed ectopic Mcl-1 as we confirmed with western blotting (Fig. 3A). This cell line was significantly less sensitive than the vector control cell line (PC-9/3M/V) to undergo apoptosis induced by the HNK and Osim combination as evidenced by reduced PARP cleavage (Fig. 3A) and limited increase in annexin V-positive cells (Fig. 3B). In agreement, the combination of HNK and Osim synergistically decreased the survival of PC-9/3M/V cells, but failed to do so in PC-9/3M/Mcl-1 cells (Fig. 3C). Collectively, these data demonstrate that enforced overexpression of ectopic Mcl-1 attenuates the ability of the HNK and Osim combination to induce apoptosis and death of Osim-resistant cells.

3.5. Bim knockout in PC-9/GA/AR cells confers resistance to the combination of HNK and osimertinib

Since in PC-9/GR/AR cells the combination of HNK and Osim clearly enhanced the levels of Bim, another critical molecule in mediating Osim-induced apoptosis in sensitive EGFR-mutant NSCLC cells (Shi et al., 2017), we determined whether enhanced Bim elevation contributes to the augmented induction of apoptosis by the HNK and Osim combination. We used the CRISPR/Cas9 technique to knock out Bim in PC-9/GR/AR cells (Fig. 3D) and then assessed their responses to treatment with HNK and Osim. Both PC-9/GR/AR/Bim-KO#2 and PC-9/GR/AR/Bim-KO#7 cells were much less sensitive than their parental PC-9/GA/AR cells to undergoing apoptosis upon cotreatment with HNK and Osim as evaluated with both PARP cleavage (Fig. 3D) and annexin V staining (Fig. 3E). Thus, Bim deficiency in PC-9/GA/AR cells confers resistance to the combination of HNK and Osim.

3.6. The combination of HNK and osimertinib enhances the growth suppression of osimertinib-resistant xenografts in nude mice

Following these in vitro studies, we then used xenograft models in nude mice to determine whether the
The combination of HNK and Osim enhances the reduction in Mcl-1 levels accompanied by augmented suppression of ERK through facilitating Mcl-1 degradation in Osim-resistant cells. (A-C) The indicated cell lines were exposed to 1 µM Osim alone, 5 µM HNK alone, or their combination for the given times (A) or 12 h (B and C) and then harvested for the preparation of whole-cell protein lysates and subsequent western blot analysis. (D) The given cell lines were treated with DMSO or 5 µM HNK plus 1 µM Osim for 8 h followed by the addition of 10 µg·mL⁻¹ CHX. At the indicated times post-CHX, the cells were harvested for the preparation of whole-cell protein lysates and subsequent western blot analysis. Band intensities were quantified by NIH Image J software, and Mcl-1 levels were presented as a percentage of levels at 0 time post-CHX treatment.
Fig. 3. Enforced overexpression of ectopic Mcl-1 or Bim-KO protects AZD9291-resistant cells from being killed by the combination of HNK and Osim as evaluated by apoptosis and cell survival. (A, B, D, and E) The given cell lines with different levels of Mcl-1 (A) or deficient in Bim (D) were exposed to 1 μM Osim alone, 10 μM HNK alone, or their combination for 48 h and then harvested for the measurement of apoptotic cells with annexin V/flow cytometry (B and E) and for the detection of PARP cleavage using western blotting (A and D). The data in B and E are means ± SDs of duplicate determinations. LE, longer exposure. (C) The indicated cell lines seeded in 96-well plates were treated with different concentrations of HNK alone, Osim alone, and their concentrations as indicated for 72 h. Cell numbers were estimated with the SRB assay. The data are means ± SDs of four replicate determinations.
combination of HNK and Osim has enhanced inhibitory effects against the growth of Osim-resistant tumors in vivo. In agreement with our observations in cell cultures described above, the combination of HNK and Osim significantly inhibited the growth of both PC-9/AR (Fig. 4A,C) and PC-9/3M (Fig. 4B,D) xenografts in comparison with vehicle control, HNK alone, or Osim alone as evaluated by measurement of both tumor size (Figs 4A,B) and weight (Fig. 4C,D). Under the tested conditions, both HNK and Osim alone did not significantly inhibit tumor growth (Fig. 4A–D). Moreover, the combination did not apparently decrease mouse body weights over the tested period (Fig. 4E,F), suggesting that the combination was well-tolerated by mice.

3.7. The HNK derivative, CAz-p, exerts similar effects in synergizing with osimertinib against osimertinib-resistant cells with increased potency

Two HNK derivatives, DEtS and CAz-p (Fig. 5A), showed greater activities than HNK in suppressing growth and inducing apoptosis of lung cancer cells as demonstrated in our previous study (Raja et al., 2008). We thus examined the effects of these derivatives in combination with Osim on the growth of Osim-resistant cells. The combination of CAz-p (e.g., 3 µM) and Osim (e.g., 1.5 µM) was clearly much more potent than either agent alone in decreasing the survival of four Osim-resistant cell lines with CIs far smaller than 1 (Fig. 5B), suggesting synergistic effects. Similar results were also generated in the colony formation assay (Fig. 5C). However, we found that DEtS combined with Osim, under the same tested conditions, did not enhance the reduction in survival of the tested Osim-resistant cell lines compared with single-agent treatment (Fig. 5D).

We further analyzed the effects of CAz-p combined with Osim on the induction of apoptosis in three Osim-resistant cell lines and found that CAz-p at 3 µM and particularly at 6 µM, when combined with Osim, was significantly more effective than either agent alone in inducing apoptosis of PC-9/AR, PC-9/GR/AR, and PC-9/3M cell lines as evaluated by measuring annexin V-positive cells (Fig. 6A) and by detecting the cleavage of caspase-3 and PARP (Fig. 6B). Moreover, the combination of CAz-p with Osim also effectively enhanced Mcl-1 reduction in these cell lines (Fig. 6C). Enforced expression of ectopic Mcl-1 in PC-9/3M cells significantly compromised the effect of the CAz-p and Osim combination on induction of apoptosis, as indicated by comparing both annexin V-positive cells (Fig. 6D) and caspase-3 and PARP cleavage (Fig. 6E). These findings suggest a critical role of Mcl-1 reduction in mediating the enhanced induction of apoptosis in Osim-resistant cells induced by the CAz-p and Osim combination.

4. Discussion

The current study provides both in vitro and in vivo evidence demonstrating that the natural product, HNK, when combined with Osim, effectively inhibits the growth of Osim-resistant cells and tumors. Hence, HNK has the potential to overcome acquired resistance to Osim. The appearance of the C797S resistant mutation is now a defined mechanism for the emergence of acquired resistance to Osim, which accounts for 20–30% of resistant cases when Osim is used as a second-line treatment (Murtuza et al., 2019). Currently, there are no effective options for the treatment of resistant tumors with triple mutations of EGFR at 19del, T790M, and C797S. The intriguing finding in our study is that the HNK and Osim combination effectively decreased the survival and induced apoptosis of both PC-9/2M cells harboring trans EGFR 19del and C797S double mutations and PC-9/3M cells carrying cis EGFR 19del, T790M, and C797S triple mutations. Another advantage of HNK is that it is a natural product used as a nutritional supplement in humans and thus has good tolerability and safety (Sarica et al., 2018). Approximately 33% of patients with EGFR-mutant NSCLC develop brain metastasis (Franchino et al., 2018), which is the main cause of mortality in this population. While Osim has improved efficacy against brain metastases (Le and Gerber, 2019), another valuable feature of HNK is that it can readily cross the blood–brain barrier and the blood–cerebrospinal fluid barrier (Wang et al., 2011; Woodbury et al., 2013), making it a suitable candidate for combining with Osim; this speculation is worthy of further investigation. Collectively, our findings warrant future evaluation of the potential of HNK to overcome Osim acquired resistance in the clinic.

The combination of HNK and Osim augmented apoptosis in the tested Osim-resistant cell lines. This enhanced induction of apoptosis is likely an important mechanism contributing to the enhanced antitumor activity of the combination of HNK and Osim against Osim-resistant cells and tumors. We have recently demonstrated that Mcl-1 reduction and Bim elevation are key mechanisms mediating Osim-induced apoptosis in sensitive EGFR-mutant NSCLC cells (Shi et al., 2017). Although enhanced Bim levels were detected only in one of three tested Osim-resistant cell lines...
Fig. 4. The combination of HNK and Osim effectively inhibits the growth of PC-9/AR and PC-9/3M xenografts in vivo. Both PC-9/AR and PC-9/3M xenografts were treated (once a day for 5 days-week\(^{-1}\)) with vehicle control, Osim, HNK, and their combination starting on the same day after grouping. Tumor sizes (A, B) were measured as indicated. Each measurement is mean ± SE (n = 6). At the end of the treatment, the mice were sacrificed to remove tumors, which were weighed (C, D). Mouse body weights were also compared (E, F). \(* P < 0.05\) at least compared with all other groups; \(** P < 0.01\) at least compared with all other groups.
Fig. 5. The HNK derivatives CAz-p and DEtS have distinctive effects in decreasing the survival and inhibiting colony formation and growth of Osim-resistant EGFR-mutant NSCLC cell lines when combined with Osim. (A) Chemical structures of HNK derivatives tested in this study. (B, D) The indicated cell lines seeded in 96-well plates were treated the next day with the given concentrations of AZD9291 alone, CAz-p or DEtS alone, or their combinations. After 72 h, cell numbers were estimated using the SRB assay. The numbers inside the graphs are CIs for the given combinations. (C) The indicated cell lines were seeded in 12-well cell culture plates. On the second day, the cells were treated with fresh medium containing DMSO, 1 µM (PC-9/GR/AR) or 2 µM (PC-9/3M) CAz-p alone, 200 nM Osim alone, and CAz-p plus Osim and the treatment was repeated every 3 days for a total of 12 days. The data are means ± SDs of four replicates (B and D) or triplicate (C) determinations. ***P < 0.001 and ****P < 0.0001 at least compared with other treatments.
Fig. 6. The combination of CAz-p and Osim enhances the induction of apoptosis in Osim-resistant EGFR-mutant NSCLC cell lines through downregulation of Mcl-1. (A,B) The indicated cell lines were exposed to DMSO, 3 or 6 µM CAz-p, 1 µM Osim, or CAz-p plus Osim for 48 h (PC-9/AR) or 72 h (PC-9/GR/AR and PC-9/3M) and then harvested for the detection of apoptosis with annexin V/flow cytometry (A) and for the detection of PARP cleavage with western blotting (B). (C) The indicated cell lines were treated with DMSO, 1 µM Osim, 6 µM CAz-p, or CAz-p plus Osim for 12 h. The proteins of interest were detected using western blotting. (D, E) The given cell lines were exposed to DMSO, 1 µM Osim, 3 µM CAz-p, or CAz-p plus Osim for 48 h (E) or 72 h (D) and harvested for measuring apoptosis with annexin V/flow cytometry (D) and for the detection of protein cleavage with western blotting (E). The data in A and D are means ± SDs of duplicate determinations. *P < 0.05, **P < 0.01, and ***P < 0.001 at least compared with other treatments.
Mcl-1 reduction than Bim elevation in mediating the induction of apoptosis by Osim and HNK combination. This is further supported by our data showing that Mcl-1 overexpression protects Osim-resistant cells from apoptosis induced by HNK and Osim co-treatment. Since Bim-KO in PC-9/GR/AP cells also protected the cells from undergoing apoptosis induced by the combination of HNK and Osim, it is fair to conclude that enhanced Bim elevation may also contribute to augmented induction of apoptosis by HNK and Osim combination at least in some Osim-resistant cell lines (e.g., PC-9/GR/AR). A recent study has shown that the natural product, bufalin, can reverse acquired resistance to Osim through induction of Ku70-mediated Mcl-1 degradation (Cao et al., 2019). Moreover, it has been shown that EGFR-mutant NSCLC cells tolerated to short-term EGFR-TKI treatment possess elevated Mcl-1 levels and can be sensitized to EGFR-TKIs by targeting Mcl-1 (Song et al., 2018). These studies together with our findings here thus highlight critical involvement of Mcl-1 in the development of acquired resistance to EGFR-TKIs including Osim and suggest a possible strategy for delaying and/or overcoming acquired resistance to Osim by co-targeting Mcl-1.

Given the dominant role of Mcl-1 suppression in mediating induction of apoptosis by Osim and HNK combination in various Osim-resistant cell lines, it is plausible to speculate that this combination may be effective against the resistance to Osim due to Bim deletion polymorphism. Hence, further study in this direction is warranted.

ERK phosphorylates Mcl-1 protein, resulting in its stabilization (Domina et al., 2004; Nifoussi et al., 2012). Osim inhibits ERK-dependent Mcl-1 phosphorylation and facilitates Mcl-1 degradation, leading to Mcl-1 reduction in sensitive EGFR-mutant NSCLC cells, as we have recently demonstrated (Shi et al., 2017). We found that the combination of HNK and Osim indeed enhanced Mcl-1 degradation. Since enhanced Mcl-1 reduction was accompanied by augmented ERK suppression (i.e., suppression of ERK phosphorylation) in the tested Osim-resistant cell lines, it is plausible to suggest that the HNK and Osim combination enhances Mcl-1 reduction or degradation likely through augmented suppression of ERK-dependent Mcl-1 phosphorylation.

One potential drawback or concern regarding the use of natural products is their relatively weak biological activities. In this study, we found that CAZ-p, a derivative of HNK, exerted similar effects as HNK did in decreasing the survival and enhancing the induction of apoptosis of several Osim-resistant cell lines including PC-9/3M, showing potential in overcoming Osim acquired resistance. CAZ-p achieved these effects at a concentration of 3 µM, which is threefold lower than that required for HNK to exert similar effects (e.g., 10 µM), indicating that CAZ-p has greater potency than HNK in overcoming acquired resistance to Osim. Therefore, it is possible to use HNK and CAZ-p as lead compounds for developing novel agents with optimized efficacy and pharmacological characteristics to overcome Osim acquired resistance.

5. Conclusions

The current study has identified HNK and its derivatives as potential candidates for overcoming acquired resistance to Osim caused by varied mechanisms such as C797S mutation. Our findings provide strong preclinical support for testing this potential strategy for overcoming acquired Osim resistance in the clinic and warrant further studies to fully understand the underlying molecular mechanisms and to develop HNK derivatives that can overcome Osim acquired resistance.

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Conflict of interest

SSR is on consulting/advisory board for AstraZeneca, BMS, Merck, Roche, Tesaro, and Amgen. TKO is on consulting/advisory board for Novartis, Celgene, Lilly, Sandoz, Abbvie, Eisai, Takeda, Bristol-Myers Squibb, MedImmune, Amgen, AstraZeneca, and Boehringer.
Ingelheim. Other people declare that they have no conflict of interest.

**Author contributions**

HZ and GQ designed and conducted experiments as well as contributed to the writing of the paper; SF contributed to the concept and paper writing; JA provided HNK and its derivatives and participated in discussion of the project; TWO and SSR contributed to the concept and paper writing; and SSY directly supervised the study, designed experiments, and wrote the paper.

**References**


