The dimer-dependent catalytic activity of RAF family kinases is revealed through characterizing their oncogenic mutants

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Abstract
Although extensively studied for three decades, the molecular mechanisms that regulate the RAF/MEK/ERK kinase cascade remain ambiguous. Recent studies identified the dimerization of RAF as a key event in the activation of this cascade. Here, we show that in-frame deletions in the β3-αC loop activate ARAF as well as BRAF and other oncogenic kinases by enforcing homodimerization. By characterizing these RAF mutants, we find that ARAF has less allosteric and catalytic activity than the other two RAF isoforms, which arises from its non-canonical APE motif. Further, these RAF mutants exhibit a strong oncogenic potential, and a differential inhibitor resistance that correlates with their dimer affinity. Using these unique mutants, we demonstrate that active RAFs, including the BRAF(V600E) mutant, phosphorylate MEK in a dimer-dependent manner. This study characterizes a special category of oncogenic kinase mutations, and elucidates the molecular basis that underlies the differential ability of RAF isoforms to stimulate MEK-ERK pathway. Further, this study reveals a unique catalytic feature of RAF family kinases that can be exploited to control their activities for cancer therapies.

Introduction
The Ras/rapidly-accelerated fibrosarcoma (RAF)/mitogen-activated protein kinase (MEK)/Extracellular signal-regulated kinase (ERK) signaling plays a crucial role in cell proliferation, survival, and differentiation [1, 2]. Aberrant activation of this kinase cascade causes developmental disorders and cancers [3–5]. Genetic alterations that hyperactivate the RAF/MEK/ERK kinase cascade exist in >40% of cancers. To target this kinase cascade for cancer therapy, both RAF inhibitors and MEK inhibitors have been developed and applied to clinical treatment [6–8]. Unfortunately, their efficacy is limited by either intrinsic or rapidly acquired resistance. Understanding the regulation of the RAF/MEK/ERK kinase cascade could help us to design strategies to circumvent this resistance and develop more effective inhibitors for cancer treatment.

The RAF kinases CRAF, BRAF, and ARAF are a core component of the RAF/MEK/ERK kinase cascade. Dimerization among RAF isoforms is a key event in triggering the RAF/MEK/ERK kinase cascade [9–18], which not only turns on the kinase activity of RAF but also facilitates the activation of MEK by RAF [19]. Mechanistic studies have shown that the two protomers play distinct roles in RAF dimers: one functions as an allosteric activator to facilitate the assembly of an active conformation in the other, which acts as a receiver to catalyze the phosphorylation of substrates [20]. Distinct molecular traits between BRAF and CRAF result in their differential ability to turn on the RAF/MEK/ERK kinase cascade by dimerization-driven
transactivation [21]. Whether ARAF can be activated by dimerization and its role in this process are unclear.

The dimerization of RAF kinase occurs in both physiological and pathological conditions, which can be induced by active Ras [13, 17], inhibitor binding [9, 11, 12, 14], gene fusions [22–25], and alternative splicings [26]. Active Ras-induced homo/hetero-dimerization of RAF kinase is responsible not only for the pathway activation by physiological agonists but also for its hyperactivation by genetic alterations upstream of RAF in carcinogenesis [13, 17, 27]. Active Ras-induced RAF dimerization can be further enhanced by RAF inhibitors, which accounts for the paradoxical effect of RAF inhibitors in cancer therapy [11, 12, 14]. It has been speculated that the RAF kinases have a close conformation in which the N-terminus docks on the carboxyl-terminal kinase domain [28]. Active Ras binds to the N-terminus of RAF kinases, which breaks their close conformation and thus facilitates their dimerization via kinase domain. On the other hand, RAF inhibitors could alter the conformation of RAF kinase domain once loaded, which not only creates a dimer-favored conformation but also relieves the inhibitory interaction between N-terminus and kinase domain [29]. The inhibitory effect of N-terminus on RAF dimerization could also be abolished by gene fusions or alternative splicing of mRNA. Chromosome translocations that lead to fusions of variable genes to the kinase domain of BRAF or CRAF have been extensively reported in cancers [22–25], while the alternative splicings that partially delete the N-terminus of BRAF(V600E) and thus enhance the dimerization of BRAF(V600E) isoforms have been shown as one of important mechanisms that underlie RAF inhibitor resistance in cancer therapy [26]. Recently, some RAF mutants (ARAF and BRAF) with in-frame deletions in the β3-αC loop have been reported as potential oncogenic drivers [30–34], although whether they are activated through enhanced dimerization remains unknown [30, 31, 33] or controversial [32, 34].

In this study, we characterized the RAF mutants with in-frame deletions in the β3-αC loop, and found that both ARAF and BRAF mutants were activated by enhanced dimerization. Further, we showed that the limited allosteric and catalytic activities of ARAF arose from its non-canonical APE motif that leads to a lower propensity of dimerization in contrast to BRAF and CRAF. Finally, we used active RAF mutants with different dimerization properties as an efficient tool to investigate whether the dimerization of RAF after activation is required for its catalytic activity and demonstrated that active RAFs including BRAF(V600E) phosphorylated MEK in a dimer-dependent manner. Our data clarifies how in-frame β3-αC loop deletions trigger RAF family kinases, reveals the molecular basis underlying the differential ability of RAF isoforms to stimulate MEK-ERK signaling, and illustrates a key step in the activation of the RAF/MEK/ERK kinase cascade.

**Results**

**Deletion of Q347A348 activates ARAF by enforcing dimerization**

By virtue of its apparent low activity and rare mutations in cancer genomes, the molecular mechanism regulating ARAF and its role in oncogenesis are ill-defined. Recently, Nelson et al. identified an active ARAF mutant with Q347A348 deletion and F351L conversion in patients with Langerhans cell histiocytosis [31]. To confirm this finding and to decipher the molecular basis of ARAF activation by this compound mutation, we expressed wild-type ARAF, and its Q347A348del mutants (ΔQA) with or without F351L mutation in 293T cell, and found that Q347A348del mutant was able to stimulate the MEK-ERK pathway independent of F351L status (Fig. 1a), suggesting a dominant role of the Q347A348 deletion in the activation of ARAF. Similar to other constitutively-active mutants of RAF kinases, a co-expression of a dominant-negative RAS mutant (N17RAS) with ARAF(ΔQA) did not affect its activity in 293T cells (Fig. 1b), indicating that ARAF (ΔQA) is a constitutively-active mutant and does not require upstream stimuli to trigger its catalytic activity. This notion was further validated by the finding that a stable expression of ARAF(ΔQA) in wild-type, BRAF−/−, and CRAF−/− fibroblasts activated the MEK-ERK pathway and transformed cells independent of endogenous RAF molecules (Figs. S1A and 1C). Moreover, the shRNA-mediated knockdown of either CRAF in BRAF−/− fibroblasts or Braf in CRAF−/− fibroblasts had no effect on the ability of ARAF(ΔQA) to stimulate downstream signaling (Figure S1B).

Dimerization of RAF molecules is critical for their activation under variable conditions [9–17]. We thus investigated whether ΔQA activates ARAF by enhancing dimerization. To do this, we carried out a complementary split luciferase assay [35] to measure the dimer affinity of ARAF and its ΔQA mutant. In this assay, the N-terminus and C-terminus of luciferase (hitherto referred to as Nluc and Cluc) were fused to ARAF and co-expressed in cells. RAF dimerization brings the Nluc and Cluc together and reconstitutes the luciferase activity. Thus, the luciferase activity correlates with the amount of dimerized ARAF. As shown in Fig. 1d, the treatment with RAF inhibitor Vemurafenib, an agonist of RAF dimerization, increased the luciferase activity of 293T cells co-expressing ARAF-Nluc and ARAF-Cluc. In contrast, the 293T cells co-expressing ARAF(ΔQA)-Nluc and ARAF(ΔQA)-Cluc showed a
constitutive luciferase activity comparable with that induced by Vemurafenib in wild-type ARAF transfectants, suggesting that ARAF(ΔQA) mutant has an elevated ability to form homodimers. The homodimerization of ARAF(ΔQA) in 293T cells can be further verified by co-immunoprecipitation assay. In contrast to its wild-type counterpart, HA-tagged ARAF(ΔQA) could be pulled down by its FLAG-tagged version when co-expressed in 293T cells although the amount is a little (Fig. 1e). Using the same method, we next evaluated the ability of ARAF(ΔQA) to heterodimerize with BRAF, CRAF, and its wild-type counterpart, and found that it barely dimerized with these molecules (Figure S1C).

It has been reported that the dimerization-driven transactivation of RAF molecules might require the interaction of the negatively charged N-terminal acidic motif (NtA motif) with the RKTRH motif in the αC-helix-β4 loop of the other protomer, and mutations that abrogate the negative charge of the NtA motif or disrupt dimer interface block this process [20]. To determine whether ARAF(ΔQA) is

**Fig. 1** The Q347A348 deletion activates ARAF by enhancing homodimerization.  

**a** ARAF(ΔQA) and ARAF(ΔQA/ F351L) have equal activity. The activity of ARAF mutants in 293T transfectants was measured by anti-phospho-ERK1/2 immunoblot. **b** The activity of ARAF(ΔQA) does not depend on upstream stimuli. ARAF(ΔQA) was coexpressed with N17Ras in 293T cells, and its activity was measured as in **a**. **c** ARAF(ΔQA) has a strong transforming ability independent of endogenous RAF molecules. Foci formation assay of immortalized fibroblasts expressing ARAF(ΔQA) was carried out as described before [36, 37]. **d, e** ARAF(ΔQA) has an elevated propensity to form homodimers. **d** The dimer affinity of ARAF(ΔQA) was measured by using complementary split luciferase assay [35]. The dimerization of wild-type ARAF induced by 10 um Vemurafenib served as a positive control (n = 5, **p < 0.001, ****p < 0.0001). **e** The dimerization of ARAF(ΔQA) was evaluated by co-immunoprecipitation assay. ARAF mutants with FLAG- or HA-tag were coexpressed in 293T cells, and immunoprecipitated by anti-FLAG beads and detected by anti-HA immunoblot. To exclude the effect of ERK1/2-mediated feedback on ARAF dimerization, all 293T transfectants in **d** and **e** were pretreated with 20 um Tramentinib for 1 h before measurements. **f** ARAF(ΔQA) is activated by dimerization-driven transactivation. Mutations that disrupt the dimer interface (R362H) or block NtA-phosphorylation (AGFF) abolish the activity of ARAF(ΔQA). The activity of ARAF mutants in 293T transfectants was measured as in a. **g, h** Homologous deletions activate BRAF in a dimer-dependent manner. g The sequence alignment of human ARAF, BRAF, and CRAF reveals conserved residues in the β3-αC loop. h, the activity of BRAF mutants in 293T transfectants was measured as in a. All images are representative of at least three independent experiments.
activated through a dimerization-driven transactivation, we mutated its NtA motif (SGYY to AGFF) or its central residue in the dimer interface (R362H), and found that both alterations impaired its activity (Fig. 1f), providing additional evidence that deletion of Q347A348 activates ARAF by enhancing homodimerization.

**Homologous deletions of two residues activate BRAF in dimer-dependent manner**

Since the Q347A348del activates ARAF by enhancing homodimerization and these two residues are conserved in the β3-αC loop across all RAF isoforms (Fig. 1g), we next asked whether a homologous deletion would activate other RAF isoforms. As shown in Fig. 1h, BRAF mutants with a deletion of either Q494L495 or Q496A497 in the β3-αC loop stimulated the MEK-ERK pathway when expressed in 293T cells, and the central RH alteration (R509H) in the dimer interface abolished their activity. This suggests that BRAF can also be activated by the β3-αC loop deletion-driven homodimerization.

**ARAF has both allosteric and catalytic activities at lower levels than those of BRAF and CRAF**

The fact that ARAF signaling through ERK was activated by Q347A348 deletion-driven homodimerization indicated that it is able to function as both allosteric activator and receiver. To confirm this, we carried out a set of RAF co-activation assays [20, 37]. In these assays, ARAF (AGFF) is a N-terminal truncated mutant (aa285-606) with a non-phosphorylatable NtA motif that functions as a receiver, whereas ARAF(DGEE/V324F) is a similar truncatant with an acidic NtA motif and a fused catalytic spine [38, 39] that mimics the inhibitor-bound status and works as a kinase-dead allosteric activator (Table S1). When co-expressed in 293T cells with allosteric activators derived from different RAF isoforms, ARAF(AGFF) was strongly activated by BRAF, immediately by CRAF, and weakly by itself through dimerization (Fig. 2a). On the other hand, ARAF(DGEE/V324F) stimulated moderately the catalytic activity of ARAF and CRAF receivers, but only slightly that of BRAF receiver when co-expressed in 293T cells (Fig. 2b, c). Taken together, these data suggest that ARAF can function as both receiver and allosteric activator, albeit less efficiently than the other two isoforms.

**The weak activity of ARAF arises from its non-canonical APE motif that decreases its dimer affinity**

Regulatory spine (R-spine) is a hallmark signature of active protein kinases that comprised of four conserved residues, namely RS1–4 [38, 39]. To explore the molecular basis underlying the weak activity of ARAF, we examined whether the R-spine-favored mutation could turn ARAF into a constitutively active kinase independent of dimerization-driven transactivation, as it has been shown for CRAF and BRAF [20, 36, 38–41]. Similar to BRAF and CRAF, an ARAF mutant combining a RS3 mutation (L358M) and a negatively charged NtA motif showed a strong activity toward MEK-ERK pathway when expressed in 293T cells (Fig. 3a). Although the activity of this mutant (ARAF, DGEE/L358M) did not depend on the activation loop (AL)-phosphorylation, it was impaired by the central RH alteration (R362H) in the dimer interface (Fig. 3b), indicating again that ARAF has different characteristics from BRAF and CRAF. By aligning the kinase domain sequences of RAF isoforms, we found that ARAF had a non-canonical APE motif whose Pro is altered into Ala (Fig. 3c). The APE motif localizes at the N-terminus of α-helix EF (αEF), and the Pro is a helix breaker that makes the N-terminus of αEF flexible. The Glu (E) next to Pro in the
The non-canonical APE motif decreases the activity of ARAF. a R-spine fusion together with acidic NtA motif fully activates ARAF. b The constitutively-active R-spine mutant of ARAF is resistant to the activation loop (AL) mutation but not to the central RH alteration in dimer interface. c ARAF has a non-canonical APE motif. The conserved APE motif is altered into AAE in ARAF, which might weaken the Glu-Arg salt bridge between APE motif and αH-αI loop. d Schematic diagram showing the Glu-Arg salt bridge in CRAF. Schematic diagram of CRAF (PDB ID: 3OMV) was generated by using PyMOL software. e The conserved APE motif restores the activity of ARAF R-spine mutant with the central RH alteration in dimer interface. f-g The conserved APE motif enhances both allosteric activator and receiver activities of ARAF. The RAF co-activation assays were carried out as in Fig. 2. h The alteration of APE motif makes BRAF R-spine mutant sensitive to the central RH alteration in dimer interface. i The alteration of APE motif or the breakage of Glu-Arg salt bridge makes CRAF R-spine mutant sensitive to the central RH alteration in dimer interface. j, k The conserved APE motif enhances the dimer affinity of ARAF mutants. The dimer affinity of ARAF mutants is measured by complementary split luciferase as in Fig. 1a (n = 5, **p < 0.001). In a, b, e and h, i the activity of RAF mutants in 293T transfectants was measured by anti-phospho-ERK1/2 immunoblot. All images are representative of at least three independent experiments. “KD” represents for “kinase domain” in the full text.
A central RH alteration in dimer interface (Fig. 3d), which has been shown to play a critical role in the regulation of protein kinase A [42]. Since Ala is a helix-favored residue, we thought that a substitution of Pro with Ala would generate a longer αEF with a more rigid N-terminus, and therefore weaken/break the Glu-Arg salt-bridge and impair kinase function. To test this hypothesis, we mutated Ala back to Pro in the APE motif of ARAF (DDEE/L358M), and found that this mutant was resistant to the central RH alteration in dimer interface as well as the homologous BRAF and CRAF mutants described in our previous studies [20, 37] (Fig. 3e). Furthermore, ARAF activator and receiver with a canonical APE motif exhibited much stronger activity than their wild-type counterparts in co-activation assays (Fig. 3f, g). On the other hand, a substitution of Pro with Ala in the APE motif of BRAF and CRAF sensitized their constitutively-active R-spine mutants, BRAF(L505M) and CRAF(DDEE/L397M), to the central RH alteration in dimer interface (Fig. 3h, i). Moreover, a breakage of the Glu-Arg salt-bridge by replacing Arg with Gin also led to the sensitivity of CRAF (DDEE/L397M) to the central RH alteration in dimer interface (Fig. 3i). Together, these data demonstrate that the different APE motifs of RAF isoforms are responsible for the differential activities observed between ARAF and the other RAF paralogs.

In order to further address how the non-canonical APE motif of ARAF dampens its functional activities, we next investigated whether it affects the dimer affinity of ARAF by using complementary split luciferase assay as described above. As shown in Fig. 3j, k, a canonical APE motif conferred the higher dimer affinity of ARAF R-spine mutants, comparing ARAF(DDEE/L358M) with BRAF (DDEE/L358M/APE), and ARAF(DDEE/L358M/R362H) with ARAF(DDEE/L358M/R362H/APE), which indicates that the non-canonical APE motif of ARAF decreases its dimer affinity. Since the dimerization plays a critical role in RAF activation, this data indicates that the lower dimer affinity arising from the non-canonical APE motif leads to the weaker activities of ARAF among RAF isoforms. In addition, the lower dimer affinity of ARAF(DDEE/L358M) and its sensitivity to the central RH alteration in dimer interface in contrast to ARAF(DDEE/L358M/APE), suggest that the active R-spine mutants of ARAF might function as a dimer to activate MEK-ERK pathway even if they do not require dimerization-driven transactivation for triggering their activity.

**Similar in-frame deletions of dimeric protein kinases exist in cancer genomes**

Dimerization-driven allostery plays a key role not only in the activation of RAF kinase but also in that of many other protein kinases [43, 44]. Besides ARAF(DQA), an oncogenic BRAF mutant with β3-αC loop deletion (ΔNVTAP) has been reported although its activation mechanism remains controversial [31–34]. We here aimed to explore all kinase mutants with similar deletions in cancer genomes and to assess the importance of these mutations in human cancers. To this end, we interrogated the ICGC (International Cancer Genome Consortium) database, the eBioportal for Cancer Genomics database, and the COSMIC (Catalogue of Somatic Mutations in Cancer) database, and summarized all kinase mutants with similar in-frame deletions of β3-αC loop, including those reported in literatures [31–34], in Table S2. Among these mutants, the EGFR exon19 del is highly prevalent and has been shown to elevate kinase activity by promoting side-to-side homodimerization [45]. Other similar kinase mutants include those of BRAF, JAK1 and ERB B2, which (except JAK1 mutants, which have not been tested) have shown elevated kinase activity (see below and Figure S2).
further demonstrate that these BRAF mutants, especially BRAF(ΔNVTAP) and BRAF(ΔMLN), are activated by enhanced homodimerization, we performed the RAF co-activation assay [20, 36, 37, 40, 41] by using their kinase-dead V471F mutants as allosteric activators. All activators tested in this study strongly stimulated the catalytic activity of CRAF receivers (Fig. 4e), and particularly activators derived from BRAF(ΔNVTAP) and BRAF(ΔMLN) could...
even trigger endogenous RAF molecules (Fig. 4e lane5 and lane3). The central R509H alteration in dimer interface was unable to prevent these two strong allosteric activators from triggering BRAF receptors (Figure S3D). Since the non-canonical APE motif had been also shown to decrease the dimer affinity in RAF molecules, we next introduced it together with the central R509H alteration into BRAF (ΔNVTAP) mutant, and found that this combined alteration completely blocked the activity of BRAF(ΔNVTAP) (Figure S3E). Taken together, our data demonstrates that all BRAF mutants with in-frame deletions of β3-αC loop are activated by enhanced homodimerization.

Unlike ARAF and BRAF, we did not find any CRAF mutants with in-frame deletions of β3-αC loop in databases. To test whether CRAF can be activated by this type of mutations, we constructed mutants homologous to those of ARAF and BRAF. When expressed in 293T cells, ΔVDPT, ΔVDPPT, and ΔVVDPPT mutants of CRAF, but not other mutants, strongly activated the MEK-ERK pathway independent of upstream stimuli (Fig. 4f and S3F&G), and exhibited differential resistance to the central RH alteration (R401H) in dimer interface (Fig. 4g). Since we had showed that BRAF was activated by mutations (ΔQA and ΔQL) homologous to ARAF(ΔQA), we hence determined whether CRAF could be triggered by mutations homologous to BRAF(ΔMLN, ΔNVTAP, and ΔNVTAPT). However, none of these alterations activated ARAF (Figure S3H).

All BRAF mutants with variable deletions of β3-αC loop have a strong transforming potential, and a robust resistance to Vemurafenib but not LY3009120 that correlates with their dimer affinity

Although the oncogenic potential and resistance to RAF inhibitor of BRAF(ΔNVTAP) has been reported recently [31, 34], whether all BRAF mutants with in-frame deletions of β3-αC loop are able to function as cancer drivers and their pharmacological characteristics are not clear. To address these questions, we first measured the oncogenic potential of these mutants by foci formation assays. As shown in Fig. 5a and S4A&B, all BRAF mutants with in-frame deletions of β3-αC loop transformed immortalized fibroblasts and induced foci formation independent of endogenous RAF molecules, suggesting that they can function as drivers to induce cancers. Further, we examined their sensitivities to the RAF inhibitor Vemurafenib, and found that all mutants exhibited a robust resistance to this drug, ranking as BRAF(ΔNVTAP) > BRAF(ΔMLN) > BRAF(ΔNVTAPT) > BRAF(ΔQA) > BRAF(V600E) (Fig. 5b, c), which correlates with their dimer affinity/stability. However, all these mutants had similar sensitivities to the RAF dimer inhibitor, LY3009120, which are comparable with that of BRAF(V600E) in A101D melanoma cell line (Fig. 5d, e).

The high dimer affinity of kinase-dead BRAF mutants with β3-αC loop deletions bypasses the requirement of active RAS to drive tumorigenesis

The inhibitor-loading has been shown to fuse the catalytic spine of RAF molecules, which can be mimicked by the Val to Phe mutation in ATP-binding pocket [20, 36–41]. Compound BRAF mutants with both β3-αC loop deletion and Val471Phe, especially BRAF(ΔNVTAP/V471F), lacked catalytic activity but were able to activate the MEK-ERK pathway through triggering wild-type RAFs (Fig. 4e). The kinase-dead BRAF(ΔNVTAP/V471F)-induced foci formation in vitro and tumor formation in vivo even in the absence of active RAS, but dependent on endogenous RAF molecules (Fig. 5f–i and S4C). Since a previous study had shown that kinase-dead RAFs cooperate with active RAS to induce tumorigenesis [12], this data suggests that the high dimer affinity of RAF mutants could bypass the requirement of active RAS to driven cancer development.

Active RAF kinases function as a dimer to phosphorylate MEK

The activation of the RAF/MEK/ERK kinase cascade is a very complex process and its underlying molecular basis is not completely understood [45–49]. In current model, RAF and MEK form a face-to-face dimer in quiescent cells. Upon stimulation, two-RAF-MEK dimers are brought together and assemble a transient tetramer through back-to-back RAF dimerization, which not only activates RAF but also facilitates subsequent MEK activation [19]. However, how active RAF activates MEK is not clear. To elucidate this process, we first tested whether the dimerization of active RAF is required for MEK activation by using BRAF mutants with in-frame deletions of β3-αC loop since these mutants have different dimer affinity/stability. As shown in Fig. 6a, those constitutively-active BRAF mutants with low dimer affinity/stability such as BRAF(ΔNVTAPT) and BRAF(ΔQA) lost their catalytic activity towards MEK in vitro when purified by immunoprecipitation, in contrast to those mutants with high dimer affinity/stability which retained catalytic activity toward MEK. We reasoned that the loss of activity of BRAF mutants in vitro arises from their dimer dissociation during purification. To test this hypothesis, we strengthened the dimers of BRAF (ΔNVTAPT) and BRAF(ΔQA), respectively, by GST fusions [50], and found that it restored their catalytic activity in vitro (Fig. 6b). This phenomenon was also seen with a homologous ARAF mutant (ΔQA) whose in vitro
catalytic activity was rescued by GST fusion (Fig. 6c). As reported before [20, 51] and shown in this study, both alterations of the central RH in dimer interface and the APE motif significantly impair but do not completely abolish the dimer formation of RAF molecules. We therefore next examined the effect of these alterations on the in vitro catalytic activity of active RAF mutants. Among three active ARAF R-spine mutants, only the one with a high
dimer affinity (see Fig. 3j, k) maintained its catalytic activity in vitro after purification (Fig. 6d), and GST fusion restored that of the other two mutants with a low dimer affinity (Fig. 6e). Similar to that of ARAF mutants, active CRAF mutants with an alteration of RH, or of APE, lost their catalytic activity in vitro, which was recovered by GST fusion-enhanced dimerization (Fig. 6f). As to active BRAF R-spine mutants, the alterations of R509H, or APE, inhibited their catalytic activity in vitro by different extends, which was also relieved by the GST fusion approach (Fig. 6g). The loss of in vitro catalytic activity of RAF mutants with low dimer affinity could be also rescued by other dimeric molecular fusions (data not shown), or partially restored with a gentle wash of PBS during purification (Figure S5), which excludes the potential artificial effect arising from GST fusion. Together, these data indicate that all RAF isoforms would function as dimers to phosphorylate MEK.

**Fig. 5** BRAF mutants with in-frame β3-αC loop deletions have a strong transforming ability and a robust but differential resistance to Vemurafenib. a BRAF mutants with in-frame β3-αC loop deletions induce foci formation when stably expressed in immortalized fibroblasts. The foci formation assay was carried out as in Fig. 1c. b, c BRAF mutants with in-frame β3-αC loop deletions exhibit a robust but differential inhibitor resistance. Stable fibroblast cells that express individual BRAF mutants with in-frame β3-αC loop deletions were treated with Vemurafenib for 4 h, and phospho-ERK1/2 was probed by immunoblot and quantified by using Image J. The graphs were generated by using GraphPad Prism 6. d-e BRAF mutants with in-frame β3-αC loop deletions have approximate sensitivities to RAF dimer inhibitor, LY309120. The drug sensitivities of BRAF mutants were measured as in b, c. The BRAF(V600E)-harboring melanoma cell line, A101D, was used as control. f-i Catalytic spine-fused BRAF mutants with in-frame β3-αC loop deletions have variable oncogenic potentials in vitro and in vivo. The oncogenic potential of BRAF mutants was measured by the foci formation assay as in a, g Xenograft tumors were generated in NOD/SCID mice from immortalized fibroblasts that express BRAF mutants as described in Materials and Methods section. h The weight of xenograft tumors from E (n = 7 for each group, **p < 0.01). i Representative images from histological section staining of xenograft tumors from E (n = 7). The MEK1DD-driven xenograft tumor served as control. All images are representative of at least three independent experiments.

**Fig. 6** Active RAF mutants phosphorylate MEK in a dimer-dependent manner. a, b BRAF mutants with in-frame β3-αC loop deletions lose their catalytic activity in vitro upon purification if they have a low dimer affinity, which is rescued by GST fusion. The band labeled with “*” in lane 2 of B represents the highly phosphorylated BRAF(KD, ΔNVTAPT). c Like BRAF(ΔQA), purified ARAF(ΔQA) loses its kinase activity in vitro, which is rescued by GST fusion. d–g GST fusion restores in vitro the kinase activity of constitutively active R-spine mutants of ARAF, CRAF, and BRAF with low dimer affinity. In a–g, all RAF mutants were expressed in 293T cells and purified by immunoprecipitation, and their activity was measured by in vitro kinase assays as described before [20, 37]. All images are representative of at least three independent experiments.
BRAF(V600E) also phosphorylates MEK through a dimer-dependent manner

Given its resistance to the central RH alteration in dimer interface and sensitivity to Vemurafenib, BRAF(V600E), a dominant mutant in RAF mutation spectrum, has been proposed to function as a monomer to activate MEK [26]. However, recent studies showed that BRAF(V600E) exists as dimers or high-order oligomers in cells [51–53]. This prompted us to determine whether, unlike other active RAF mutants, BRAF(V600E) truly phosphorylates MEK in a dimer-independent manner. Since BRAF(V600E) has an enhanced propensity to form dimers [51–53], the central RH alteration (R509H) in dimer interface is unable to dissociate its dimers completely. Hence, our approach was to replace one protomer in the dimer of BRAF(V600E) with one dysfunctional BRAF mutant (loss of both catalytic and allosteric activities), and measured the catalytic activity of BRAF(V600E) in heterodimers. To generate such a dysfunctional BRAF mutant, we mutated the residues of the kinase-dead BRAF(ΔNVTAP/V471F) that mediate the heterodimerization of BRAF with MEK [19] (Fig. 7a). Since BRAF utilizes two different groups of residues to bind MEK and RAF [19], the introduction of these mutations would not influence RAF dimerization. Unlike its prototype, the mutant, BRAF(ΔNVTAP/V471F/R462E/ΔI loop, heretofore referred to as BRAF(ΔNVTAP/V471F), had no allosteric ability to trigger endogenous RAF molecules when expressed in 293T cells (Fig. 7b). Moreover, BRAF(V600E) bound to BRAF(ΔNVTAP/V471F)* had little catalytic activity in vitro, in contrast to that bound to BRAF(ΔNVTAP/V471F) when purified from 293T co-transfectants (Fig. 7c). This suggests that a BRAF (V600E) molecule needs a partner able to hold MEK in order to phosphorylate it. To further confirm this finding, we introduced BRAF(ΔNVTAP/V471F)* into BRAF(V600E-dependent melanoma cell lines by lentiviral transductions and found that its expression downregulated phospho-ERK1/2 and inhibited cell growth in vitro and xenograft tumor growth in vivo (Fig. 7d–h).

As shown above, the APE motif of RAF kinases regulates their dimerization, likely through affecting the Glu-Arg salt bridge between the APE motif and the αH-αI loop, and the alteration of APE motif into non-canonical AAE dramatically decreases RAF dimer affinity/stability. Therefore, we next determined whether an alteration of APE motif into AAE would impair the catalytic activity of BRAF (V600E) in vitro, as which occurs in ARAF R-spine mutants. As shown in Fig. 7i, j and S6A, although the AAE variant of BRAF(V600E) had comparable ability with its prototype to phosphorylate MEK and thus turn on ERK signaling when expressed in 293T cells, it lost its catalytic activity upon purification by immunoprecipitation, which was restored by GST fusion. Moreover, the combined alterations of AAE and R509H completely blocked the activity of BRAF(V600E) even in vivo, which was also recovered by GST fusion (Fig. 7i). To directly confirm that the discrepant catalytic activity among BRAF(V600E), BRAF(V600E/AAE), and BRAF(V600E/AAE/R509H), arises from their different dimer affinity/stability, we completed a co-immunoprecipitation assay with a gentle wash of PBS as in Figure S5, and found that these BRAF variants had quite different ability to form dimers as BRAF (V600E) » BRAF(V600E/AAE) » BRAF(V600E/AAE/R509H) (Figure S6B). Altogether, these data demonstrate that like other RAF molecules, BRAF(V600E) also functions as a dimer to activate MEK.

Discussion

The dimerization of RAF kinase not only plays a critical role in the activation of the RAF/MEK/ERK kinase cascade [9–17], but also contributes to drug resistance in cancer therapy [11, 12, 14, 26]. Previous studies have shown that the RAF dimerization could be improved by active RAS, inhibitor binding, gene fusions or alternative splicing [9, 11–14, 17, 22–26, 54]. However, whether it can be achieved by other ways remains unknown. Recently, some oncogenic RAF mutants with β3-αC loop deletions have been reported by several groups [31–34], and both dimer-dependent and -independent models have been suggested to explain how this type of mutations activates RAF [32, 34]. Chen et al. characterized BRAF(ΔLNVTAP) that resembles BRAF(ΔLNVTAPT) in cancer genomes, and found that its activity was blocked by the central R509H alteration in dimer interface [32]. On the other hand, Foster et al. showed that the other cancer-related mutant, BRAF(ΔNVTAP) was resistant to the same alteration [34]. Both groups failed to understand the discrepancy among these RAF mutants. In this study, we systemically characterize all RAF mutants with β3-αC loop deletions, and provide solid evidence that this type of mutations activates RAFs through improving homodimerization, which clarifies the controversial between these two groups.

Among RAF isoforms, BRAF and CRAF have been shown to function as both catalysis-competitive kinases and allosteric activators [21], while ARAF has a bare activity and been thought as a scaffold [55, 56]. By characterizing the ARAF mutant with a β3-αC loop deletion (ΔQA), we demonstrate that like the other two paralogs, ARAF has both catalytic and allosteric activities though less. Further, we elucidate that the weak activity of ARAF arises from its non-canonical APE motif that decreases its dimer propensity likely by weakening the Glu-Arg salt bridge between the APE motif and the αI-αH loop.
explain why ARAF has less ability to stimulate downstream MEK-ERK signaling.

In-frame deletions of β3-αC loop activate not only RAFs but also other dimeric protein kinases by enhancing homodimerization. Remarkably, we found this type of mutation in BRAF, ERBB2, JAK2, and EGFR in cancer genomes, suggesting that the dimerization works as a principal mechanism to regulate the activity of these
The dimer-dependent catalytic activity of RAF family kinases is revealed through characterizing their mutual dependence on active RAS proteins. Recent studies have shown that the catalytic activity of RAF kinases is significantly enhanced when they are present in a dimeric state. This dimerization is crucial for the activation of downstream pathways, particularly the MEK-ERK cascade.

**Materials and methods**

**Biochemicals**

Antibodies used in this study included: anti-phosphoERK1/2 (#9102), anti-phosphoMEK1/2 (#9150), and anti-RAF (αS/C loop loop exhibit the elevated albeit differential dimer affi- nity, which is reflected by their different resistance to the central R509H alteration in dimer interface. The strong dimer affinity of BRAF(NVTAP) bypasses the require- ment of active RAS for its kinase-dead version to activate endogenous RAF and transform cells, suggesting that the dimerization of RAF is a dynamic equilibrium in cells and active RAS shifts this equilibrium in the favor of dimers.

A strong dimerization is required for transactivation of RAF kinases, whereas a weak dimerization is still indispensable for their catalytic function. The previous conclusion that active RAF kinase may function as a monomeric enzyme is mainly based on the data from the central RH alteration in dimer interface of RAF molecules. Recent studies revealed that this alteration was not able to completely dissociate RAF dimers although it blocked the dimerization-driven transactivation of RAF molecules [20, 51]. Here, we found that this mutation does not block the dimerization-driven transactivation of RAF mutants with high dimer affinity, such as BRAF(NVTAP) and BRAF(AML2), which calls for a re-evaluation of previous data resulting from the central RH alteration in dimer interface. Constitutively active RAF mutants including BRAF(V600E) activate MEK in a dimer-dependent manner even if they do not require a dimerization to trigger their activity, implying that RAF kinase serves as both an enzyme and a MEK-docking platform in its catalysis process. This is further supported by the kinetic-dense BRAF mutant with high dimer affinity is not able to activate downstream MEK-ERK pathway through wild-type RAF molecules if it cannot bind MEK (Fig. 7a, b). These findings suggest that the MEK-docking platform function of RAF kinase can be used as a target to develop novel therapeutic inhibitors against both RAS- and RAF-driven cancers. In current cancer therapies, the efficacy of RAF and MEK inhibitors has been severely limited by intrinsic and acquired resistances arising from the paradoxical activation or re-activation of RAF/MEK/ERK kinase cascade [57]. Such a new class of docking inhibitor would cover all type of cancers driven by hyperactive RAS/RAF/MEK/ERK signaling, and thus potentially have a greater and longer efficacy. Our study sheds a light on the treatment of RAS/RAF-driven cancers and has important clinical implications.

**Plasmids and cell lines**

cDNAs encoding proteins in this study were purchased from Addgene or synthesized by Integrated DNA Technologies. All mutations were generated by PCR and tagged with either FLG or HA or His, and cloned into vectors by Gibson assembly. pCDNA3.1(+) vector (Invitrogen) was used for
transient expression; viral vectors (Clontech) for stable expression; and pET-28a (Novagen) for bacterial expression. Wild-type, B-RAF\(^{−/−}\) and CRAF\(^{−/−}\) MEFs were generated in previous study [58, 59]. Melanoma cell lines: MeWo, A101D, Mel-624 were obtained from ATCC.

**Protein expression and purification**

6xhis-tagged MEK1 (K97A) and 6xhis-tagged ERK2 (K52A) were expressed in BL21(DE3) strains and purified by using Nickel column (Qiagen) and following our previous protocol [60].

**Cell culture, transfection, and transduction**

All cell lines were maintained in DMEM medium with 10% FBS (Hyclone). Cell transfection were carried out by using the Biotool transfection reagent and following the manufacturer’s protocol. To generate stable cell lines that express RAF or MEK1 mutants, viruses were prepared and applied to infect target cells according to our previous studies [20, 36, 37, 60, 61]. Infected cells were sorted by FACS or selected by using antibiotics.

**Immunoprecipitation, in vitro kinase assay, and Western blotting analysis**

Immunoprecipitations were performed as described previously [20, 36, 37]. Briefly, whole-cell lysates were mixed with either anti-HA (E6779), or anti-FLAG beads (A2220) (Sigma), rotated in cold room for 60 min, and washed three times with RIPA buffer. For in vitro kinase assays, the immunoprecipitants were washed once with kinase reaction buffer (25 mM HEPES, 10 mM MgCl\(_2\), 0.5 mM Na\(_3\)VO\(_4\), 0.5 mM DTT, pH 7.4), then incubated with 20 μl kinase reaction mixture (2 ug substrate and 100 mM ATP in 20 μl kinase reaction buffer) per sample at room temperature for 10 min. Kinase reaction was stopped by adding 5 μl per sample 5XLaemmli sample buffer. Immunoblotting was carried out as described before [20, 36, 37, 60, 61].

**Foci formation assay**

Immortalized MEFs infected with retroviruses encoding target proteins were plated at 5 × 10\(^5\) cells per 60 mm dish, and fed every other day. Twelve days later, cells were fixed with 2% formaldehyde and stained with Giemsa solution (Sigma).

**Complementary split luciferase assay**

A total of 293T transfectants that express different pairs of Nluc- and Cluc-fused RAF proteins were plated in 24-wellKrystal black image plates at the the seeding density of 2 × 10\(^5\) per well. Twentyfour hour later, D-luciferin (0.2 mg/ml) with or without Vemurafenib (10 μM) was added to the culture, and the incubation was allowed for 30 min before the luciferase signals were measured by Promega GloMax\textsuperscript{®}-Multi Detection System.

**Animal studies**

For xenograft experiments, female NOD/SCID mice (6–8 weeks) were injected with 3 × 10\(^6\) cells per mice in 1:1 matrigel (Corning). Tumor volumes were monitored by digital calipers twice a week and calculated using the formula: volume = (width)\(^2\) × length/2. At the experiment endpoint, mice were killed and tumors were collected for ex vivo analysis and subsequent histology. All operations were approved by the Animal Ethics Committee of NCCS.

**Immunohistochemistry staining**

Tumors were fixed in 10% buffered formalin overnight and embedded according to standard procedures. Tumor sections were cut to 4 μm thickness, mounted on glass slides, and air-dried at room temperature. After antigen retrieval, tumor sections were stained with antibodies and then with hematoxylin. Images of tumor sections were taken with a bright light microscope at 10×.

**Statistical analysis**

All statistical analysis in this study was performed using GraphPad InStat (GraphPad Software, CA, USA). Statistic significance was determined by two-tailed Student’s \(t\)-test in animal studies and error bars represent s.d. to show variance between samples in each group, or by one-sample \(t\)-test in other experiments and error bars represent s.d. to show variance between independent experiments.

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**Author contributions**

JY and JH designed the study; JY and JH searched databases/literatures for kinase mutations in cancer genomes; MB prepared experimental materials; JY, WHN, YW, HX, JJY, and AL performed immunohistology analysis; PL, MW, and JH supervised all experiments and interpreted experimental data; JY wrote the manuscript; MB and MW revised manuscript; and all authors commented and approved the manuscript.
Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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