Activation of K-RAS by co-mutation of codons 19 and 20 is transforming

Adam Naguib1, Catherine H Wilson2, David J Adams2, Mark J Arends3*

Abstract

The K-RAS oncogene is widely mutated in human cancers. Activating mutations in K-RAS give rise to constitutive signalling through the MAPK/ERK and PI3K/AKT pathways promoting increased cell division, reduced apoptosis and transformation. The majority of activating mutations in K-RAS are located in codons 12 and 13. In a human colorectal cancer we identified a novel K-RAS co-mutation that altered codons 19 and 20 resulting in transitions at both codons (L19F/T20A) in the same allele. Using focus forming transformation assays in vitro, we showed that co-mutation of L19F/T20A in K-RAS demonstrated intermediate transforming ability that was greater than that of individual L19F and T20A mutants, but less than that of G12D and G12V K-RAS mutants. This demonstrated the synergistic effects of co-mutation of codons 19 and 20 and illustrated that co-mutation of these codons is functionally significant.

Findings

Mutations in RAS family genes occur in approximately 20-30% of all human cancers, with mutations in the K-RAS gene comprising ~80% of these mutations [1]. K-RAS mutations have been documented in the majority of human cancer types with pancreatic (~90% of these cancers) and colorectal (~40%) cancers demonstrating the highest incidence of mutations in this gene [2,3]. K-RAS codons 12 and 13 are the most common sites of oncogenic activation with over 90% of documented mutations being found in these codons [4]. Amino acid alterations at these codons, which encode amino acids adjacent to the GDP/GTP binding pocket, reduce or abolish GTPase activity of K-RAS after GAP binding and lock the protein in an active, GTP-bound state [5]. Codons 12 and 13 in wildtype K-RAS both encode glycine residues. The incorporation of other amino acids, most commonly aspartate and valine at codon 12 and aspartate at codon 13 [6], brings about projection of larger amino acid side chains into the GDP/GTP binding pocket of the protein, interfering with the geometry of the transition state in which GTP hydrolysis is catalysed [7]. Mutations in codons 61 and 146 have also been described to be oncogenic in K-RAS, although mutations at these positions occur at a much lower prevalences (<5% of total K-RAS mutations) than codon 12 and 13 mutations [4]. Activation of RAS has several effects on rodent cells in vitro, including the establishment of a transformed phenotype with anchorage independent growth in soft agar, transformed focus formation, as well as tumour formation following injection into animals [8-10]. Our previous mouse model studies have shown that induced expression of mutant K-RAS accelerates intestinal adenoma formation in vivo on both a mutant Apc background and a Msh2-null background, increases proliferation, modulates apoptosis and alters gene expression patterns [11-15]. In this study, we identify a novel double mutation outside of codons 12 and 13, recoding K-RAS codon 19 from leucine to phenylalanine (L19F) and codon 20 from threonine to alanine (T20A), in a colorectal cancer and we demonstrate that it causes transformation in vitro and is therefore a rare, but functionally significant co-mutation of K-RAS.

Sequencing of exons 1 and 2 of K-RAS in a panel 186 colorectal adenocarcinoma samples, obtained as part of the European Prospective Investigation into Cancer and Nutrition (EPIC) Norfolk study cohort, identified 41 cancers harbouring mutations in K-RAS [16]. Forty of these mutations were located in codons 12 or 13, however, one sample demonstrated the presence of double mutant peaks, one at the third position of codon 19 (G > T giving rise to a leucine to phenylalanine (L19F) amino acid change) and the other at the first position of codon 20
individual PCR amplicons of $K$-RAS exon 1 harbouring the L19F/T20A mutations. Analysis of sequencing traces from individual bacterial clones identified solely double mutant (L19F/T20A) alleles (Figure 1), thus confirming the presence of both mutations on the same allele. In order to confirm the somatic nature of both elements of this co-mutation, non-cancerous DNA from the blood of the same individual was sequenced. Blood samples were obtained upon the volunteer’s enrolment into the initial EPIC study. Following tumour development, both blood and tumour tissue were made available for analysis and processed for DNA extraction [16]. Sequencing of genomic DNA and sequencing methodology has been described in detail elsewhere [16].

Missense mutations of $K$-RAS codon 20 in human cancers have not been previously reported, however, mutations giving rise to phenylalanine incorporation into $K$-RAS codon 19 have been described previously in seven individual human colorectal cancers, a single human lung adenocarcinoma sample and a single lymphoblastic leukaemia [17-20]. No observations of a double mutation giving rise to both L19F and T20A amino acid alterations have been described previously. As such, the observation in our study of double mutant peaks giving rise to L19F/T20A missense co-mutations at codons 19 and 20 of $K$-RAS describes a previously unreported change in this gene.

A series of full length human $K$-RAS isoform B cDNA sequences were designed and synthesised containing either G12V, G12D, L19F, T20A, combined L19F/T20A or wildtype sequences and these were cloned into pBABE-puro expression vector plasmids for transfection into mouse NIH3T3 cells for focus forming assays (see Additional File 1). Transfection of $K$-RAS cDNA constructs containing individual L19F or T20A mutations did not show significantly increased transformation above that of the control cells, which were treated with either transfection reagents but no cDNA or transfection reagents and wildtype $K$-RAS cDNA. However, cells transfected with constructs containing the double mutation L19F/T20A in $K$-RAS cDNA demonstrated significantly increased focus formation above that of wildtype $K$-RAS cDNA transfected control cells (Mann-Whitney U test: $P = 0.03$; Figures 2 and 3). Additionally, the $K$-RAS L19F/T20A co-mutation was shown to form fewer foci than cells transfected with either G12D or G12V mutated $K$-RAS cDNA ($P = 0.02$ and 0.04 respectively). These data demonstrated that this L19F/T20A co-mutation of $K$-RAS confers oncogenic activation capable of inducing transformed focus formation to a greater degree than single mutations at either codons 19 or 20 or wildtype $K$-RAS cDNA, but to a lesser degree than codon 12 mutants of $K$-RAS.

It is not surprising that the T20A substitution alone was not sufficiently oncogenic to cause focus formation as this amino acid alteration has never been found previously in human cancers. However, the L19F substitution has been documented, albeit in a limited number of studies, and may be expected to have some transforming potential. One study described analysis of L19F in the *C. elegans* RAS homologue *let-60*. *C. elegans* carrying this mutation showed a temperature-sensitive multivulval phenotype. In a mammalian system, at body temperature, H-RAS (L19F) protein had a reduced rate of GTP hydrolysis relative to wildtype H-RAS, suggesting that H-RAS L19F conferred an increased level of activation [21]. However, transfection of NIH3T3 cells with human H-RAS with the incorporated L19F mutation failed to demonstrate increased focus formation above that of controls, a similar observation to that made here using L19F $K$-RAS. A second report, however, describes
L19F as causing increased cell proliferation, anchorage-independent growth, increased tumourigenicity in nude mice and elevated levels of RAS-GTP [17]. Additionally, a recent analysis describing K-RAS mutations outside of codons 12 and 13 also described mildly increased focus formation by L19F mutants [22]. In this report the L19F K-RAS mutation was described to induce formation of ~5 foci whereas positive controls (G12V and G12D) were observed to develop ~80 - 90 foci. These data are not mutually exclusive with our observations, as the mean focus count observed here with the G12D mutant was 35 colonies per plate, thus a mild focus forming ability of the L19F mutation may not have been detected. The observed differences between the number of colonies formed in the study by Smith and colleagues [22] and those in our own assays may be due to variations in protocols used, such as the different expression vectors used in the two analyses giving rise to different levels of transcription following transfection into NIH3T3 cells and different tissue culture plating conditions.

Our identification and analysis of the L19F/T20A double mutation in K-RAS is the first description of this genetic change, which confers a greater transforming ability than individual mutations in these codons. The rare observed frequency of this double amino acid change giving rise to oncogenic K-RAS in human cancers, compared with that of the individual L19F amino acid change, despite its increased oncogenic potential, may be due to the lower likelihood of 2 base changes occurring at these positions in the same allele during cancer development.

To assess the potential structural effects of the L19F/T20A amino acid changes we modelled their side chains into a shortened wildtype K-RAS protein ribbon structure bound to a GTP analogue (Figure 4a). Codons 19 and 20 (shown in white in Figure 4b) are in close proximity to the GDP/GTP binding pocket of the protein. Modelling of the mutant phenylalanine (codon 19) and alanine (codon 20) side chains into the wildtype protein predicted two features. First, the removal of the leucine side chain at codon 19 and its replacement with that of phenylalanine resulted in projection of the bulky, ringed side chain of the newly encoded amino acid into the main body of the protein and this may be predicted to change the shape of the GDP/GTP pocket by steric interference. Second, removal of the threonine side...
chain at codon 20, with loss of its OH group and its replacement with alanine that has a smaller aliphatic side chain may be predicted to affect the ionic interactions between nearby amino acid side chain groups (Figure 4c & 4d). The presence of changes at both codons 19 and 20, demonstrating a higher transforming activity than individual mutations in these codons, suggests that shifting of the protein conformation due to steric interference together with disruption of ionic interactions within the K-RAS protein, may be responsible for generating a protein configuration capable of providing the observed transforming activity, by interfering with the geometry of the GDP/GTP binding pocket reducing GTP hydrolysis. Further structural studies are required to test this proposed mechanism.

K-RAS mutations affecting codons 15 and 22 have also been reported in human colorectal cancers [23,24]. Although the transforming ability of these specific codon changes is yet to be confirmed, the observation of sequence changes in human cancers affecting codons 15 and 22, as well as at co-mutation of codons 19 and 20, as described here, strongly suggests that alteration of this region in the K-RAS protein provides a selective growth advantage for cells which contributes to neoplastic transformation.

Additional material

Additional file 1: Experimental methods and statistical testing
Additional File 1 contains a detailed description of all experimental methods described in this manuscript. Furthermore, complete colony counts are available for all experiments in tabulated format.

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Author details
1Medical Research Council Dunn Human Nutrition Unit, Wellcome Trust/MRC Building, Cambridge, CB2 0XY, UK. 2Wellcome Trust Sanger Institute, Wellcome Trust Genome Campus, Hinxton, Cambridge, CB10 1SA, UK. 3Department of Pathology, University of Cambridge, Addenbrooke’s Hospital, Cambridge, UK, CB2 0QQ, UK.

Authors’ contributions
All authors contributed to the study design, analyses and manuscript preparation. All authors read and approved the final manuscript.
Competing interests
The authors declare that they have no competing interests.

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