Post-irradiation somatic mutation and clonal stabilisation time in the human colon

F Campbell, G T Williams, M A C Appleton, M F Dixon, M Harris, E D Williams

Abstract

Background—Colorectal crypts are clonal units in which somatic mutation of marker genes in stem cells leads to crypt restricted phenotypic conversion initially involving part of the crypt, later the whole crypt. Studies in mice show that the time taken for the great majority of mutated crypts to be completely converted, the clonal stabilisation time, is four weeks in the colon and 21 weeks in the ileum. Differences in the clonal stabilisation time between tissues and species are thought to reflect differences in stem cell organisation and crypt kinetics.

Aim—To study the clonal stabilisation time in the human colorectum.

Methods—Stem cell mutation can lead to crypt restricted loss of O-acetylation of sialomucins in subjects heterozygous for O-acetyltransferase gene activity. mPAS histochemistry was used to visualise and quantify crypts partially or wholly involved by the mutant phenotype in 21 informative cases who had undergone colectomy up to 34 years after radiotherapy.

Results—Radiotherapy was followed by a considerable increase in the discordant crypt frequency that remained significantly increased for many years. The proportion of discordant crypts showing partial involvement was initially high but fell to normal levels about 12 months after irradiation.

Conclusions—Crypts wholly involved by a mutant phenotype are stable and persistent while partially involved crypts are transient. The clonal stabilisation time is approximately one year in the human colon compared with four weeks in the mouse. The most likely reason for this is a difference in the number of stem cells in a crypt stem cell niche, although differences in stem cell cycle time and crypt fission may also contribute. These findings are of relevance to colorectal gene therapy and carcinogenesis in stem cell systems.

Keywords: colon, clonality, stem cell, mutation, radiation, cell kinetics.

The intestinal epithelium is a stem cell tissue, which is organised into structurally coherent clonal units with each unit maintained by one or more stem cells; the clonal units have been identified morphologically as the crypts of Lieberkühn. The hierarchy of cells involved in the maintenance of clonal units has been investigated using markers such as tritiated thymidine or bromodeoxyuridine but the fact that the nuclear activity is diluted with each successive division following a pulse label and that the great majority of S-phase cells that take up the label are not stem cells make it difficult to determine accurately the organisation and kinetics of stem cells, particularly as these cannot be identified separately.

Phenotypic changes caused by mutation are capable of giving rise to a permanent change in the clonal unit if the mutation occurs in a stem cell. We have previously shown that mutagen induced loss of glucose-6-phosphate dehydrogenase (G6PD) activity visualised by histochemistry can be used to demonstrate such a clonal phenotypic change in widely scattered crypts in the mouse colon. Time course studies allowed the evolution of crypt colonisation by the mutant phenotype to be followed. Five days after a single administration of mutagen most of the discordant crypts were partially involved by cells expressing low levels of the G6PD mutated phenotype. The frequency of these diminished over the four weeks after mutagen administration as the number of discordant crypts in which the whole clonal unit was occupied by mutant cells increased. The frequency of wholly discordant crypts reached stability at about the same time as the frequency of partially involved crypts fell to very low levels. Similar findings have been made in the mouse small intestine either using G6PD or a technique that detects mutations in mice heterozygous for a gene controlling lectin binding, but here the time taken to reach the plateau in the frequency of wholly discordant crypts was up to five times longer than in the colon. We consider it likely that this surprising difference between small and large intestines is due to tissue variation in the number and organisation of stem cells located within a crypt stem cell niche, although differences in stem cell kinetics or crypt fission would also contribute. Whether this or other models of crypt organisation are used, the time taken to reach the plateau represents the time taken for the progeny of one stem cell to populate the whole clonal unit in nearly all mutated crypts. This may be termed the clonal stabilisation time.

We have also demonstrated a monogenically inherited polymorphism for the histochemical staining of human colorectal mucosa by a technique (mPAS) that discriminates...
O-acetylated (mPAS negative) from non-
O-acetylated (mPAS positive) epithelial sia-
lo-mucoproteins. Somatic mutation of the high
acetylator allele in crypt stem cells in
heterozygous subjects followed by crypt
colonisation by the mutant progeny leads to
conversion of the crypt phenotype from mPAS
negative to mPAS positive and we have used
this to identify and quantify somatic mutations
in colectomy specimens. Radiation given
one month before colectomy for carcinoma led
to a considerable increase in frequency of
mPAS positive discordant crypts, but the
increase was very largely confined to crypts
showing partial involvement by the mutant
phenotype, and no information is available on
the clonal stabilisation time in human colon.
Unlike G6PD histochemistry, the mPAS
technique can be used on archival formalin
fixed tissues. We have therefore used mPAS
staining to study mutation in colectomy
specimens from patients who had radiation to
the pelvic area at varying times up to 34 years
before colectomy. The results suggest that the
time taken for the progeny of one stem cell to
replace all the cells of a colonic crypt in
humans is approximately 12 months.

Methods
Colectomy specimens were available for study
from 38 patients who had received prior
therapeutic radiation. The age of the patients
at radiation ranged from 31 to 77 years, the age
at operation from 33 to 79 years, and the
elapsed time between radiation and colectomy
from four months to 34 years. There were 31
women. The radiation was given as treatment
to carcinoma of the cervix (18 cases), bladder
(4 cases), anorectum (3 cases), endometrium
(2 cases), prostate (2 cases) or other tumours
(9 cases). Colonic resection was undertaken for
colorectal carcinoma in three cases, a sigmoid
colon adenoma in one, symptoms attributed to
radiation proctocolitis in 33 cases, and for
obstruction in one. The radiation dose given
ranged widely, from 4000 cGy of fractionated
external beam radiation in a patient with
carcinoma of the rectum to a combination of
4000 cGy of fractionated external beam
radiation and 4800 cGy of intracavitary
radiation to a patient with carcinoma of the
cervix. The relationship of the part of colon
sampled to the field irradiated could not be
determined accurately, particularly in the cases
where intracavitary radiation was given, as the
tissue dose falls rapidly with distance from the
source.

One hundred and twenty consecutive large
bowel resection specimens for sporadic
primary colorectal cancer were used for
comparison. The patients were aged 28 to 94
years at the time of surgery and there were 57
women. None of these patients had received
pre-operative radiotherapy or cancer chemo-
therapy.

The tissues studied consisted of formalin
fixed paraffin wax embedded blocks of
resection margins taken during routine
histological examination for diagnostic pur-
poses. In non-irradiated colons these were
invariably normal macroscopically and micro-
scopically whereas some blocks from irradiated
colons showed minor histological changes of
radiation colitis. However blocks showing
active inflammation, ulceration or severe
distortion of the mucosal architecture were
excluded. One 5 µm section from each case
was stained with the mPAS technique11 to
discover if the phenotype was homozygous for
low O-acetyl transferase activity (showing
mPAS positive staining of all crypts). All other
cases were largely or completely negative on
staining with mPAS and further multiple
mPAS stained step sections from blocks of each
case were obtained such that at least
10 000 crypt profiles (the minimum required
for confident assessment of mutated crypt fre-
quencies12) were examined for each subject.8 10
These step sections were cut at 80 µm intervals
to obtain the necessary size of sample while
keeping the chances of counting the same crypt
(normal or mutated) in adjacent sections at a
low level. The total sample of crypt profiles
examined in each case was determined by
counting manually all of the profiles in one
central step section (using a hand held tally)
and multiplying this figure by the number of
step sections examined (usually 10–20 per
case).

In all cases with mPAS positive crypts in an
mPAS negative background the number of
discordant crypts was counted, and each
recorded as partially or wholly replaced by the
mutant phenotype. Results were expressed as
the frequency of either wholly involved or
partially involved crypts ×10−4. Statistical
comparisons between patient groups were
made using the Mann-Whitney U test.

Results
Thirty eight cases where colectomy had been
performed four months or more after
completion of a course of abdominal radiotherapy
were available for study. Thirty were
unselected cases, of which four showed
uniformly low levels of O-acetylation (mPAS
positive), nine were excluded because less than
10 000 crypts were available for study, and four
showed uniformly high levels of O-acetylation
(mPAS negative). This left 13 cases with the
heterozygous phenotype, showing high levels
of O-acetylation and scattered crypts in which
the mPAS negative phenotype was completely
or partially replaced by cells that had lost
O-acetylation (mPAS negative with scattered
positive crypts). These cases were suitable for
quantitation of somatic mutation frequency. A
further eight irradiated cases with discordant
mPAS positive crypts in an mPAS negative
background were derived from other studies,
giving a total of 21 informative cases.

The 120 specimens with sporadic colorectal
cancer untreated by radiation or chemotherapy
included 15 uniformly mPAS positive, 52
uniformly mPAS negative, and 53 predomi-
nantly mPAS negative with scattered mPAS
positive crypts. More than 10 000 crypts were
available for assessment in all of these cases.
The frequency of wholly involved discordant crypts in the 21 informative irradiated cases was very variable, ranging from $4.3 \times 10^{-4}$ to $12.3 \times 10^{-4}$, as were the frequencies of partially involved crypts which ranged from 0 to $39.1 \times 10^{-4}$. The overall mean (SD) and median for all 21 informative cases was 37.7 (35.1) $\times 10^{-4}$ (median 26.3 $\times 10^{-4}$) for wholly affected crypts, 7.8 (10.4) $\times 10^{-4}$ (median 3.4 $\times 10^{-4}$) for partially involved crypts, and 45.5 (39.8) $\times 10^{-4}$ (median 33.8 $\times 10^{-4}$) for all affected crypts. These figures compare with 10.9 (11.7) $\times 10^{-4}$ (median 7.5 $\times 10^{-4}$) for wholly affected crypts, 1.0 (1.0) $\times 10^{-4}$ (median 0.9 $\times 10^{-4}$) for partially affected crypts, and 11.8 (12.1) $\times 10^{-4}$ (median 7.7 $\times 10^{-4}$) for all affected crypts from the 53 informative control colorectal cancer specimens that had not received prior radiation.

The frequencies of wholly involved, partially involved and all discordant crypts show a statistically significant increase in the irradiated compared with the non-irradiated patients (Mann-Whitney U, p < 0.0001 for each comparison).

To evaluate the results in relation to the time period after radiation, the 21 irradiated cases were divided into three groups of approximately equal size, 4–12, 13–24, and >25 months after radiotherapy. Although the tissue radiation dose could not be determined accurately the range of radiation given was approximately the same in each of the groups. The results are presented in the Table, along with those from the non-irradiated cases and a previously published study of seven similarly aged informative subjects given 4000 cGy of fractionated external radiation one month before operation for colorectal cancer. Because the data were not normally distributed, the results are shown as medians and ranges. The frequency of wholly discordant crypts rises significantly (p < 0.003) 4–12 months after radiation, and remains raised thereafter without changing significantly. The frequency of partially involved crypts is also significantly increased 4–12 months after radiation (p < 0.002) and then falls with time. However, it does not return to the low frequency seen in non-irradiated cases, the median value of $2.1 \times 10^{-4}$ at >25 months being significantly higher (p < 0.01) than $0.9 \times 10^{-4}$ observed in the non-irradiated group.

A wide variation in the frequencies of discordant crypts between cases in each of the three groups of irradiated cases studied would be expected because it was not possible to determine the spatial relation between the exact part of the colon available for study and the radiation field. We have therefore set out the results as the proportion of all discordant crypts that are due to partially involved crypts.

The Table shows that this proportion, with a median value of 6% in non-irradiated cases, was considerably increased at 56% one month after radiation, and fell progressively with time thereafter to reach 16% at 4–12 months, 13% at 13–24 months, and 10% at >25 months. The values at one month and 4–12 months post-radiation are significantly increased compared with non-irradiated cases (p < 0.0001 and p < 0.03 respectively).

**Discussion**

This study shows that after exposure to therapeutic levels of irradiation the human colon shows the same pattern of changes in discordant (mutated) crypts as we have seen in mouse colon after exposure to a chemical carcinogen, but at a very different time scale. It confirms our previous findings that the frequency of mutated crypts identified by mPAS histochimistry in informative cases rises significantly as a consequence of exposure to radiation. At one month the increase is largely due to crypts that are partially involved by the mutant phenotype; with increasing time after irradiation wholly involved crypts predominate and the frequency of partially involved crypts declines. The frequency of wholly affected crypts appears to reach a plateau after about 12 months, remaining increased for many years thereafter. The overall findings support previous conclusions that wholly affected crypts are persistent and result from fixed mutations of the O-acetylttransferase gene in

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**Frequencies of discordant (mutated) crypts**

<table>
<thead>
<tr>
<th>Time since radiation (months)</th>
<th>Number of informative cases</th>
<th>Median age in years at time of colon resection (range)</th>
<th>Radiation dose range (cGy)</th>
<th>Median frequency of discordant crypts $\times 10^{-4}$ (range)</th>
<th>Median % of discordant crypts that were partially involved (range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 month</td>
<td>53</td>
<td>73 (49 to 94)</td>
<td>none</td>
<td>7.5± (0 to 53.4)</td>
<td>6% (0 to 50*)</td>
</tr>
<tr>
<td>4 to 12 months (11 months)</td>
<td>7</td>
<td>64 (31 to 73)</td>
<td>4000 external</td>
<td>6.95 (0 to 17-9)</td>
<td>5% (26-9 to 61.9)</td>
</tr>
<tr>
<td>13 to 24 months (19 months)</td>
<td>5</td>
<td>54 (33 to 70)</td>
<td>5000 internal+</td>
<td>3.12 (4.3 to 14.1)</td>
<td>13 (2.7 to 16.7)</td>
</tr>
<tr>
<td>25 to 408 months (53 months)</td>
<td>9</td>
<td>65 (46 to 79)</td>
<td>4800 internal+</td>
<td>2.14 (5-8 to 128)</td>
<td>10 (0 to 45-5)</td>
</tr>
</tbody>
</table>

*excluding one case with only a single partially involved crypt.

†data from reference 10.

Statistically significant differences (Mann-Whitney U): 
- 54-irradiated v 4–12 months, 13–24 months and >25 months: p < 0.03;
- 54-irradiated v 1 month, 4–12 months, 13–24 months and >25 months: p = 0.04;
- 54-irradiated v 4–12 months, 13–24 months and >25 months: p = 0.03;
- 54-irradiated v 1 month and 4–12 months: p = 0.03.
long lived stem cells while partially affected
crypts are transient, resulting either from relatively recent stem cell mutations whose
progeny have not yet colonised the whole
crypt, or post-stem cell mutations in the pro-
liferative compartment of the crypt.10

A particular point of interest is the decline with time after radiation in the proportion of
all discordant crypts that is composed of partially involved crypts (Table), a ratio that
animal studies suggest is independent of the
tissue dose.4 7 In a previous study of mutation
in mouse colon using a different marker (the
X-linked enzyme G6PD) we also found a fall
with time in this proportion after a single dose of
mutagen, either ethylnitrosourea or di-
methylhydrazine.4 This change in the relative
proportions of wholly involved crypts and
partially involved crypts could be explained on
the basis that there is a single crypt stem cell
with a long cycle time, partially involved crypts
resulting from mutations in relatively long lived
non-stem cells or in stem cells repopulating the
crypts. A more plausible explanation is that the
crypts are maintained by a stem cell niche in
which there are two or more stem cells per
crypt and the loss of cells from the niche after
cell division is at least partly random.1

Mutation in one stem cell will then maintain
a mixed phenotype crypt until either the
mutated stem cell is lost or the mutated stem
cell line replaces the non-mutated stem cells.
In the mouse colon, after a single mutagen
dose, the proportion of all discordant crypts
formed by mixed phenotype crypts returns to
normal in about four weeks.4 5 In the mouse
small intestine the time is between 125 and
212 8 weeks, depending on which experimental
model is used. We have termed this the clonal
stabilisation time – the time taken for
phenotype stability to be achieved within
clonal units (in this case colonic crypts) after
a single exposure to a mutagen. Our current
findings suggest that in the human colon it is
of the order of 12 months. It is unlikely to be
much less than one year because the median
percentage of partially involved crypts is
significantly higher than the non-irradiated
cases at both one month and 4–12 months
post-irradiation and remains higher at 13–24
months after irradiation (though the latter
difference does not reach statistical signifi-
cance). Moreover the time since irradiation is
not randomly distributed in the 4–12 month
group, but is skewed towards the upper limit
of the time range with a median of 11 months.

The very long clonal stabilisation time in the
human of about 12 months compared with the
four weeks in mouse colon could result from
a number of factors including a larger number
of stem cells in the crypt stem cell niche in
humans or a longer stem cell cycle time. It
could also be influenced by an alteration
affecting the probability that the loss of a cell
from the niche occurs at random. This raises
the possibility that a mutation affecting a
growth control gene or one involved in cell
signalling could influence this probability,
giving a mechanism that could reduce the
chance of potentially carcinogenic mutations
persisting in stem cells. Whether this
mechanism exists remains to be proved, but
the kinetics of the crypt stem cell are clearly
relevant to the process of colorectal neoplasia.

Several other issues require consideration,
including the persistence of an increased
frequency of partial mutated crypts over many
years, the unusually high proportion of the
group of irradiated patients who were
apparently heterozygotes for O-acetyltrans-
ferase activity, and mechanisms other than the
crypt niche hypothesis, which could explain
the change in the relative proportion of partially
involved crypts with time.

The crypt niche model for three stem cells
predicts an asymptotic fall in the number and
proportion of partially involved crypts with
time after mutagen exposure.4 If a niche
contains a greater number of stem cells,
sufficient for one to be surrounded by other
stem cells, the shape of the niche and the
position of the mutated stem cell within it will
influence the time taken for the crypt to reach
phenotypic stability. The long continued
increase in the absolute frequency of partially
affected crypts is therefore explicable on a stem
cell niche model. We have considered the
possibility that partially involved crypts
continue to be generated because of a rise in
mutation rate that persists for years after
exposure to high dose radiation – a number of
in vitro mutagenicity studies, recently reviewed
by Kronenberg,13 have suggested that x
radiation not only produces somatic mutation
from direct DNA damage to radiated cells but
also induces a persistent destabilisation of the
gene predisposing to chromosomal instab-
bility and a permanent increased suscepti-
bility to mutation in the surviving progeny.
However, because of the similarity of the total
mutated crypt frequencies at 4–12, 13–24, and
>25 months after irradiation it is unlikely that
continuing new stem cell mutations resulting
from such a late irradiation effect make more
than a small contribution to the raised partially
involved crypt count.

The frequency of mutated crypts cannot
unfortunately be related on an individual case
basis to the tissue exposure to radiation in this
study. The uncertainty is due to the rapid fall in
dose with distance from the intracavitary source
used in many cases, and the lack of knowledge
of the precise anatomical location of the portion
of colon from which the blocks studied were
derived. It is therefore probable that in some or
all of the three irradiated cases with the lowest
level of discordant crypts (<9 × 10−3) the colonic
mucosa assessed was outside the radiation field.
An alternative explanation that must be
considered is that some of these cases were
homozygous rather than heterozygous for a high
level of O-acetyl transferase activity. Of the 30
unselected cases, four showed diffuse mPAS-
positivity, consistent with homozygosity for low
O-acetyltransferase activity. This proportion is
similar to that found in previous studies.8
However, in this study the proportion of cases
with discordant crypts was higher than expected
both from previous studies, and from application
of the Hardy-Weinberg law (although the
Post-irradiation somatic mutation in the human colon

difference is not statistically significant by the χ² test, p > 0.1. This may result from chance, or it could be the consequence of mutation in homozygotes. The highest total mutation rate observed in a single case was 144.4 × 10⁻⁴. The mutation frequency in homozygotes where two hits are required to produce loss of function, is predicted to be the square of that in heterozygotes, where one hit only is needed. A mutation leading to a discordant crypt frequency of 144.4 × 10⁻⁴ in a heterozygote would therefore be expected to give a frequency of 2·1 × 10⁻⁴ in a homozygote. This is a detectable level, and would be higher if, as is probably the case, the mutations were clustered in one part of the colon because of a steep gradient in tissue dose. The relatively high proportion of apparent heterozygotes may well result from chance, but we cannot exclude the possibility that some of the cases with a low count are due to mutations in homozygotes exposed to very high levels of radiation.

It has recently been suggested that crypts preparing to undergo fission increase their stem cell number, that these crypts are particularly liable to stem cell mutation, and that the conversion of a partially involved to a wholly involved crypt takes place at the time of crypt fission. In this model crypts would be maintained by more than one stem cell, but the control of asymmetry could be intrinsic to the stem cell. Crypt fission clearly is a mechanism that could contribute to the conversion of a partially mutated crypt to a wholly mutated crypt and a phenotypically normal crypt, although if more than two stem cells are present per crypt several rounds of crypt fission would probably be required to give a uniform phenotype. We have rarely observed a wholly involved crypt adjacent to a partially involved crypt, which could have arisen by fission of a partially involved crypt with a minimum of three stem cells at the time of crypt fission. Crypt fission rates decline with age so that if this was an important factor in the susceptibility to mutation and in clearance of partially involved crypts, the stem cell mutation rate would be expected to fall with age and the clearance rate to lengthen. In fact we find experimentally that 30 days after a single dose of mutagen the colonic somatic mutation frequency induced in 27 month old mice is higher than in 3 month old animals but the ratio of wholly to partially involved crypts is very similar in both age groups (unpublished findings). For this and other reasons we think it unlikely that crypt fission plays more than a minor part in the clearance of partially involved crypts.

We regard the clonal stabilisation time as the time taken for virtually all genetically altered cells either to be lost from a clone or to replace all the cells of a clone. In the colon we regard the crypt as the clonal unit, even though we consider it likely to be maintained by a stem cell niche, and we have shown that the crypt stabilisation time in humans is about one year. Knowledge of the mechanisms that control this time and lead to species and tissue differences is relevant to theories of carcinogenesis. Knowledge of the time itself is relevant to gene therapy, as it is the interval that must pass before permanent levels of introduced genes can be guaranteed.


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