ISOENZYMES OF HEXOKINASE, 6-PHOSPHOGLUCONATE DEHYDROGENASE, PHOSPHOGLUCOMUTASE AND LACTATE DEHYDROGENASE IN UTERINE CANCER

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Summary.—Electrophoresis of cytosol prepared from normal and malignant tissue samples of uterine cervix and endometrium revealed interesting differences which may be relevant to the characteristic alterations in glucose metabolism associated with tumour development. Hexokinase II was detected in 30% of the cancer material from both sources, but in none of the samples of normal cervix. A duplet band of 6-phosphogluconate dehydrogenase was seen in the majority of the cancer samples but in no sample of normal cervix; it appeared to be partly due to ageing of the sample, and is not phenotypically related to the malignant process. Analysis of genetic variance for phosphoglucomutase at the PGM₁ locus revealed a highly significant excess of the PGM₁-1 phenotype in patients with cancer of the endometrium, which may reflect susceptibility to endometrial cancer in patients with this phenotype. At the PGM₂ locus, samples of malignant cervix were deficient in “Band f” compared with normal cervix samples, all of which showed this band. Conversely, gene products of the PGM₃ locus were found in most samples of malignant cervix and a small minority of normal cervix samples. Compared with the isomorphic distribution of lactate dehydrogenase enzymes in normal uterine tissue, cancers showed a shift towards either a more anodal or a more cathodal pattern. The former may be associated with tumours enjoying a good oxygen supply, and the latter with tumours which, because of their depth or poor blood supply have to function under less aerobic conditions.

We have previously reported a significant increase in activity of enzymes concerned with glucose metabolism in cancers of the human cervix uteri compared with normal cervical epithelium (Marshall et al., 1978a). This increase was noted for enzymes of the direct oxidative pathway and for those of the glycolytic pathway. Kinetic studies demonstrated differences in behaviour of aldolase and pyruvate kinase of normal and malignant cervical epithelium, with respect to substrate affinity and response to certain activators and inhibitors, and suggested that cervical cancers may differ from normal cervical epithelium not merely in having higher total enzyme activity but also in having different molecular forms of these enzymes (Marshall et al., 1978b). We were unable to obtain satisfactory results with zone electrophoresis to answer this question for these two enzymes, but were stimulated to apply these techniques to other enzymes in the hope that such efforts might throw some light on the regulation of these metabolic pathways in human tumours, and might reveal phenotypic differences between normal and malignant cells. This paper reports our findings with 4 enzymes with which differences in the

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electrophoretic pattern given by normal and malignant cervical epithelium were
seen with relatively high frequency.

MATERIALS AND METHODS

General.—Samples of normal and malignant tissue from the cervix and body of the uterus
were collected, homogenized and centrifuged to prepare 105,000g supernatant fractions for
assay of enzyme activities, as described in our earlier report (Marshall et al., 1978a), which
also lists the sources of the reagents which were used in the present work. Protein con-
centration was determined by the method of Lowry and haemoglobin according to Drabkin
& Austin (1932). The pH of all solutions was adjusted at 20°C. Starch gels were prepared
at a concentration of 12-5% (w/v) in buffer, and horizontal electrophoresis was performed
at a constant voltage of 200V for 18 h at 5°C Gel slices were developed at 37°C for 2-3 h in
the case of hexokinase, 6-phosphogluconate dehydrogenase and phosphoglucomutase. Lac-
tate dehydrogenase isoenzymes were separated on a slab of 5-5%, polyacrylamide gel prepared
from the following solutions: A, 48 ml n HCl, 36-3 g Tris, 0-23 ml TEMED, water to 100 ml
and pH adjusted to 8-9; B, 30 g acrylamide, 0-8 g N,N-methylene bis-acrylamide and 100
ml water; C, 140 g ammonium persulphate and 100 ml water. Each gel slab was prepared
by mixing the following: 2 ml solution A, 2 ml solution B, 8 ml solution C, and 5-09 ml
water to yield a final gel concentration of 5-5%. Electrophoresis was performed in the
vertical position at a constant voltage of 200 V for 3 h at 5°C, after which the gels were
sliced and stained for 15 min at 37°C.

With all enzymes, normal and malignant samples were included on each gel slab, and
were diluted as necessary so that none was greater or less than 25% of the mean activity
for all samples applied. Preliminary experiments carefully defined the activity limits and gel-staining times so that no isoenzyme band reached maximum intensity under the conditions used (unless otherwise stated). Strict proportionality was thus main-
tained between each sample, and between the isoenzyme components within each sample.

Specific conditions.—(a) Hexokinase (EC 2.7.1.1; ATP: D-hexose 6-phosphotrans-
ferase). These were based on the method of Brewer & Sing (1970). The gel buffer comprised
0-021m Tris, 0-02m boric acid and 0-68m
EDTA, disodium salt, at pH 8-6. The tank buffer was 0-21m Tris, 0-15m boric acid and
6mm EDTA, disodium salt, adjusted to pH 8-0. The stain was 50mm Tris/HCl, pH
7-68; 8mm MgCl2; 2mm glucose; 4mm ATP; 0-2mm NADP; 0-24mM nitroblue tetrazolium
(NBT); 0-33mM phenazine methosulphate (PMS); and glucose 6-phosphate dehydrogen-
ase (G6PD), 0-33 u/ml.

(b) 6-Phosphogluconate dehydrogenase (EC 1.1.1.44; 6-phospho-D-gluconate-NAD+ oxido-
ductase, decarboxylating). These were based on the method described above for hexokinase (Brewer & Sing, 1970). The gel and tank buffers were as for the previous enzyme, except that NADP was added to the cathode compartment to give a final concentration of 0-02mM to improve resolution and yield bands of higher intensity. The stain comprised 100mM Tris/HCl, pH 7-8;
10mM MgCl2; 0-2mM NADP; 0-6mM 6-phosphogluconate; 0-33mM PMS; and 0-24mM
NBT.

c) Phosphoglucomutase (EC 2.7.5.1; α-D-glucose-1, 6-biphosphate:α-D-glucose-1-
phosphate phosphotransferase). These were based on the method of Spencer et al.
(1964). The reservoir buffer was 0-1m Tris; 0-1m maleic acid; 0-01m EDTA, disodium salt;
and 0-01m MgCl2; and the pH was adjusted to 7-6 with 5m NaOH. The gel buffer was a 10-fold dilution of the reservoir buffer. The stain comprised 100mM Tris/HCl, pH 7-3;
5mM MgCl2; 0-2mM NADP; 5mM glucose 1-phosphate (containing 1% glucose
1,6-diphosphate); 0-24mM NBT; 0-33mM PMS; and G6PD, 0-33 u/ml.

d) Lactate dehydrogenase (EC 1.1.1.27; L-lactate: NAD+ oxidoreductase). These
were based on the method of Dietz & Lubrano (1967). The reservoir buffer consisted of
5mM tris and 38mM glycine adjusted to pH 8-3. The stain was 50mM Tris/HCl, pH 8-0;
500mM sodium lactate; 2mM NAD; 0-24mM NBT; and 0-33mM PMS.

RESULTS

Hexokinase

Katzen & Schimke (1965) separated 4 zones of hexokinase activity from rat
tissues, designated HKI–HKIV in order of increasing anodal mobility. HKI, II and III could be detected when the staining solution contained 5×10^{-4} m
glucose, whereas HKIV, found in liver, required 0·1M glucose concentration in the stain. HKIII, in contrast to HKI and II, was inhibited by glucose concentrations in the staining solution greater than about 10\(^{-2}\)M, a result confirmed for human hexokinase isoenzymes by Rogers et al. (1975). In the present work no additional bands of activity were demonstrated in 11 normal and malignant specimens stained with solutions containing 0·1M glucose as opposed to 2mM glucose and, therefore, because of the danger of inhibiting HKIII if present, a concentration of 2mM glucose in the staining solution was routinely used for development of hexokinase isoenzymes.

Only samples with activity greater than 100 mu/ml proved sufficiently active for our detection system, of which 19 were from normal cervix epithelium, 19 were from malignant cervix and 8 were from malignant endometrium.

All showed a low mobility zone with a duplet structure presumably corresponding to the duplet zone of HKI described by Rogers et al. (1975). Many malignant samples, both of cervix and endometrium, showed another more mobile band corres-

![Image of starch gel electrophoresis](image.png)

**Fig. 1—Electrophoresis of hexokinase isoenzymes on starch gel.** Origin (O) marks point of application of 5 malignant samples, all of which show duplet HKI isoenzyme, and Nos 1, 2 and 4 (from left) also demonstrate a zone of HKII activity towards the anode (+).
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PICTURE 2—Electrophoresis of 6-phosphogluconate dehydrogenase on starch gel. Origin (O) marks application of 4 malignant samples, 3 of which (Nos 1, 2 and 4 from left) demonstrate the duplet band towards the anode (+).

Corresponding with HKII (Fig. 1), HKII was not detected in any of 19 samples of normal cervix, but was present in 6/19 samples of cervical cancer and 3/8 samples of endometrial cancer. Normal endometrium had insufficient activity to enable detection of hexokinase isoenzymes. No hexokinase isoenzymes other than I and II were observed in the material examined.

6-Phosphogluconate dehydrogenase

Of 25 normal cervix epithelium samples, all showed a single band of uniform mobility towards the anode. Malignant samples revealed a slightly slower band in addition to that of normal specimens (Fig. 2) in 22/32 samples. This additional band was seen in 19/25 samples of cervical cancer and in 3/7 samples of endometrial cancer. These specimens had been stored at −20°C for up to 2 months. To check these findings, analyses were performed on a number of samples within 1 week of preparing the supernatant. When this was
done, 10 normal and 23 malignant samples gave single bands, suggesting some structural change had taken place on storage, with reduction of net charge, increase in size, or both.

To accelerate the process of duplet formation, normal and malignant samples were incubated at 37°C, alone, and in the presence of 0.2 mm NADP or 0.5 mm mercaptoethanol before electrophoresis. After 24 h at 37°C, 3/5 malignant samples showed the appearance of a duplet, whereas 6 normal samples were unaltered. NADP and mercaptoethanol had no further effect. No correlation between haemoglobin content and the presence of double bands was found. It is unlikely that erythrocytes are the source of the second band.

### Phosphoglucomutase

This enzyme is thought to be coded in 3 separate loci, designated PGM1, PGM2 and PGM3 (Harris et al., 1968). PGM1 contains 3 common genetic variants noted by Spencer et al. (1964) which are transmitted by autosomal inheritance although, according to a more recent report (Bark et al., 1976), further common phenotypes may

#### Table 1.—Distribution of PGM1 phenotypes in normal and malignant uterine tissue samples

| PGM1 Phenotype | Normal Cervix | Normal Endometrium | Cancer Cervix | Cancer Endometrium |
|----------------|---------------|-------------------|---------------|-------------------|
| 1              | 15            | 9                 | 17            | 33                |
| 2-1            | 16            | 3                 | 11            | 4                 |
| 2              | 1             | 1                 | 2             | 1                 |

Fig. 3.—Electrophoresis of phosphoglucomutase on starch gel, with diagrammatic representation of phenotypes at each of the 3 gene loci for this enzyme. The samples applied at the origin (0) demonstrate for the PGM1 locus the following phenotypes 2, 1, 2-1, 2-1, and 1, respectively. The gene products at the other loci are not clearly resolved on this gel.
be detected by isoelectric focusing. Each homozygous phenotype presents after electrophoresis as an intense slow band and a more mobile band of lower intensity. The common heterozygotic phenotype of PGM1 appears as 4 bands with mobilities and relative intensities of a mixture of the 2 homozygotes—that is, no hybridization occurs. Fig. 3 illustrates diagrammatically the appearance of the phenotypes and shows a typical electrophoretic separation. All 3 phenotypes occurred in normal and malignant cervix and endometrium, and their frequency is reported in Table I.

Fig. 4.—Electrophoresis of phosphoglucomutase on starch gel. Origin (0) marks application of 6 malignant samples. Prolongation of running and staining periods reveals 3 bands in first (from left) and 2 bands in the remainder at the PGM3 locus. All samples show an "e" band and the first sample both "e" and "f" bands at the PGM3 locus.
Harris et al. (1968) reported the distribution of PGM phenotypes in a random sample of the British population: 58% exhibited PGM$_1^1$, 36% showed PGM$_1^{2-1}$ and the remaining 6% had PGM$_1^2$. The combined normal samples of cervix and endometrium in the present study showed 53% PGM$_1^1$, 42% PGM$_1^{2-1}$ and 4% PGM$_1^2$, in good agreement with Harris et al. (1968). Malignant cervix samples did not vary significantly from these frequencies, but malignant endometrium showed a very high frequency of the PGM$_1^1$ phenotype ($\chi^2 = 11.02; P < 0.01$).

Genetic variants of the second locus, PGM$_2$, are rare. The usual form seen in erythrocytes consists of 3 bands decreasing in intensity with increasing mobility. In the uterine tissues studied only 2 bands were detected, the second (Band f) being absent in 13/18 malignant cervix samples, but present in all 18 normal cervix samples.

PGM$_3$ variants are common but could only be detected in samples with a high activity, as they accounted for a small proportion of total phosphoglucomutase. Even then, staining beyond the usual 3 h needed for optimal resolution of PGM$_1$ bands was necessary. Because of this and their high mobility, it was difficult to

### Table II

**Lactate dehydrogenase isoenzyme patterns in normal and malignant uterine tissue samples**

| Pattern       | Normal | Cancer |
|---------------|--------|--------|
|               | Cervix | Endometrium | Cervix | Endometrium |
| LD$_1$-shift  | 4      | 5      | 10     | 9      |
| Symmetrical   | 32     | 3      | 7      | 4      |
| LD$_2$-shift  | 10     | 2      | 20     | 3      |

**Fig. 5.**—Electrophoresis of lactate dehydrogenase on polyacrylamide gel. Origin (O) marks beginning of running gel, with isoenzymes numbered LD$_5$ to LD$_1$ with increasing mobility towards the anode (+). The forward horizontal line shows the albumin–bromophenol-blue marker. Mixed normal and malignant samples were applied showing the following patterns: 1 and 3, LD$_1$- shift; 2 and 4, LD$_2$ shift; remainder symmetrical.
distinguish them while simultaneously resolving PGM\(_1\) variants. PGM\(_3\) was represented by 2–3 fast bands detected in 15/18 malignant cervix samples and 2/18 normal cervix samples (Fig. 4). Because of their faintness, we could not assess the frequency of variants at the PGM\(_3\) locus.

**Lactate dehydrogenase**

Three patterns were distinguished in the samples studied: a symmetrical distribution about LD\(_3\); a shift towards LD\(_1\); and a shift towards LD\(_5\). Table II demonstrates that, in general, normal cervix epithelium had a symmetrical pattern of LDH isoenzymes, whereas malignant cervix showed a shift towards either LD\(_1\) or, more commonly, LD\(_5\) (Fig. 5). The pattern of endometrial cancer was comparable to that of normal endometrium, both showing LD\(_1\)-dominance.

**DISCUSSION**

**Hexokinase**

Hexokinase isoenzyme shifts involving an increase in HKII have been described in chemically-induced tumours of rat liver, mammary gland, and kidney (Shatton et al., 1969), in experimental rat hepatomas (Sato et al., 1969), in human hepatomas (Balinsky et al., 1973; Hammond & Balinsky, 1978) and in all of 53 primary human tumours investigated (Kamel & Schwarzfischer, 1975).

The appearance of HKII in malignant uterine tissues is in agreement with the observations of Kikuchi et al. (1972) but, whereas they found HKII in all 10 samples of malignant cervix examined, only one third of the uterine tumours in our series showed HKII. On the other hand, Kikuchi et al. (1972) detected HKII in 3/10 samples of normal cervix, which may indicate contamination of their material with erythrocytes, which contain large amounts of HKII (Rogers et al., 1975), or greater sensitivity of their detection system. In the present work, erythrocytes were removed by dissecting out blood clots and washing thoroughly with water. No mention of these precautions was made by Kikuchi et al. (1972). Haemoglobin estimations revealed very low levels in all but 4 of our samples demonstrating HKII, and the presence of the latter cannot be attributed to erythrocytes.

Since HKI and HKII differ in molecular weight, thermostability and appearance during foetal development, it has been suggested that they are determined by separate gene loci (Rogers et al., 1975). Appearance of HKII in tumours would therefore represent reprogramming of protein synthesis. It has been suggested that this may be necessary for progression to neoplasia after the initiation event, and would permit a higher rate of glucose phosphorylation (Kamel & Schwarzfischer, 1975). Recent work supports the suggestion that increased HKII levels augment glucose utilization (Bernstein, 1977). Our failure to detect HKII in the majority of tumours precludes this change being an obligatory event in neoplasia. This conclusion is supported by a recent paper describing the presence of HKII in only 13/27 human glial tumours (Bennett et al., 1978).

**6-Phosphogluconate dehydrogenase**

Latner (1967) reported a single band of PGDH in cervical carcinomas after starch-gel electrophoresis. The mobility was variable from tumour to tumour, but inequalities of mobility were abolished by adding excess NADP. He offered 3 explanations for this variability of the isoenzyme: (a) gene mutation; (b) ageing of cells in the population; and (c) variable NADP content of the tumours. Differences in electrophoretic mobility of 6-phosphogluconate dehydrogenase from malignant melanoma and normal skin have been described (Prasad et al., 1974) Tumour tissue occasionally showed a slower-moving band which was also present in muscle. The two bands differed in their binding capacity for NADP. During storage, many malignant samples developed a slower band of 6-phosphogluconate dehydrogenase after electrophoresis. Pos-
sible reactions resulting in loss of mobility are: loss of a cofactor, deamidation, desialylation, oxidation of sulphhydril groups, and conformational changes (Épstein and Schechter, 1968). Preincubation of normal and malignant samples with NADP or mercaptoethanol, however, evoked no change in mobility, whereas incubation at 37°C for 24 h resulted in duplet bands in 3/5 malignant samples. This reflects a difference in the microenvironment between the normal and the malignant state; perhaps the redox potential or the level of protein catabolism is the relevant factor.

**Phosphoglucomutase**

Assuming that the phenotypes observed for this enzyme in the uterus are characteristic of the individual genotype, the association between the PGM1\(^1\) phenotype and women with endometrial carcinoma may reflect association between the disease and some genetically determined aetiological factor. Women with PGM1\(^2\) may be less susceptible. A comparable situation has been demonstrated by Cassimos et al. (1973), who showed an association between glucose-6-phosphate dehydrogenase deficiency and a low incidence of cancer.

In the tissues studied, 2 instead of the 3 bands of PGM2 seen in erythrocyte haemolysates were detected. This may represent genuine tissue variation or a feature of the electrophoretic technique. The second (Band f) was not detected in 13/18 malignant cervix samples, whereas all 18 normal cervix samples showed it. The most intense band (e), which was always present, may be the primary gene product and Band f may arise by subsequent modification; absence of Band f might suggest inactivity of the modifying agent in most cervical cancers.

Two or three fast bands representing the PGM3 locus were detected in 15/18 malignant cervix samples and 2/18 normal samples. This difference in detection of PGM3 bands presumably reflects an activity increase of enzymes of this locus due to malignant change. It is not due to higher total phosphoglucomutase activity of cancer samples *per se* since, as with other enzymes, the amount loaded on to the gel was quite constant. The significance of such an increase is uncertain. Although differences in specificity, affinity and molecular weight are known between the various PGM gene products (Harris et al., 1968), the function of PGM3 is enigmatic. PGM1 is thought to be mainly involved with glucose-1-phosphate (G1P) as substrate, whereas PGM2 uses ribose-1-phosphate (R1P) more efficiently. PGM3 has a very high Km (10\(^{-2}\)M) for G1P and does not use R1P significantly.

**Lactate dehydrogenase**

Okabe et al. (1968) described a symmetrical pattern for the lactate dehydrogenase isoenzymes of normal cervix. Many workers (Turner, 1964; Sutcliffe & Emery, 1968; Latner et al., 1966; Fottrell et al., 1974) reported an increased LD\(_5\) in samples of cervical carcinoma. Thus previous work is in agreement with the present findings, except for a study by Langvad & Pedersen (1969), who reported no significant increase in the ratio of LD\(_4\) to LD\(_2\) in 54 patients with carcinoma of the cervix. Their electrophoretic method did not allow visualization of LD\(_5\), and there is difficulty in observing an LD\(_2\)→LD\(_4\) shift compared with an LD\(_1\)→LD\(_5\) shift. This finding of increased LD\(_5\) content is in line with extensive studies of other human tumours (Goldman et al., 1964; Nissen & Bohn, 1965) and is postulated to permit increased rates of glycolysis, since LD\(_5\) is less susceptible to inhibition by pyruvate (Latner & Skillen, 1967).

The surprising observation of an LD\(_1\) shift in about a quarter of the samples of carcinoma of the cervix has not been previously reported, although this phenomenon has been observed in other tumours (Henderson et al., 1974; Prasad et al., 1974). The proportion of H-subunits in human hepatomas seemed to be slightly greater than in normal or host liver (Hammond & Balinsky, 1978). Haemoglobin determinations revealed a some-
what high content in 3/19 tumour samples with LD₁ shift; haemoglobin concentrations in the same range were present in 7 tumour samples demonstrating LD₅ shift. It is therefore improbable that erythrocyte contamination can account for the observed instances of LD₁ shift in the malignant material.

Lactate dehydrogenase (LDH) isoenzyme patterns may change in tumours according to physiological stimuli rather than as a consequence of reprogramming of protein synthesis necessary for neoplastic growth. Oxygen tension, hormonal changes and virus interaction are reported to cause differential production of LDH subunits (Clark & Yochim, 1969; Prasad et al., 1972). Perhaps the LDH isoenzyme pattern reflects the oxygen tension in a tumour. Superficial tumours directly accessible to oxygen may show an LD₁ shift, whereas deep tumours with poor vascularization may show an LD₅ shift.

No such correlation was attempted in this work, but would be worth trying in future.

Significant differences in LDH isoenzyme patterns were not observed in endometrial carcinoma compared with normal endometrium. Fotrell et al. (1974) reported 70–76% M-subunits in 3 cases of carcinoma of human endometrium compared with 11% in normal endometrium samples during the proliferative phase of the menstrual cycle and 32% during the secretory phase. It is difficult to assess the significance of this work because of the small number of tumours examined, and because the variance of results in the control material was not indicated.

CONCLUSION

Differences in the isoenzyme patterns for the 4 enzymes examined have been demonstrated between normal and malignant tissues of the human uterus. In the main, these changes are qualitative rather than absolute, and they did not show a significant degree of correlation in individual tumours or in the tumour population as a whole. No cancer-specific isoenzymic forms were elucidated. The closest to this objective was the failure of expression of the PGM₃ locus in normal cervix epithelium and the frequent occurrence of a duplet form of 6-phosphogluconate dehydrogenase in malignant cervix which was never seen in normal tissue. Further work is needed to elucidate the chemical basis for this phenomenon, and the genetic basis for the other differences, which may be due to changes in the cell population making up the biopsy sample. The relatively small number of samples did not encourage a detailed analysis of the relationship between the observed phenomena in cancers and the type of tumour, its sensitivity to therapy, its degree of invasiveness, and whether these phenomena have any prognostic significance. These goals are worthy of future study.

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