It is clearly established that hypertension is a major factor in the genesis of arterial disease in adult life (Kannel and Dawber, 1974). Severe hypertension in childhood is a relatively uncommon but important disorder, for in at least 80% of cases it is secondary to some underlying, often treatable, condition (Loggie, 1971; Kaufman et al., 1972). Furthermore, it is increasingly being realized that essential hypertension may have its origins in childhood (Londe et al., 1971). The renin-angiotensin system plays an important causative or perpetuating role in many forms of hypertension; for this reason a review of its status in children and its relevance to childhood hypertension is appropriate.

**Renin angiotensin system (Fig.)**

Renin is a proteolytic enzyme produced by the juxtaglomerular cells of the afferent arterioles in response to a variety of stimuli. The most important of these is a decrease in renal arterial perfusion pressure, but changes in renal tubular fluid sodium concentration and sympathetic nervous activity also play a part. The renin is released into the circulation and acts on an α2-globulin, renin substrate, generating the physiologically inactive decapetide, angiotensin I. In the plasma and on passage through the lungs, angiotensin I is converted to the active pressor octapeptide, angiotensin II, which causes arteriolar vasoconstriction and hence an increase in total peripheral resistance. Angiotensin II also has a less well understood direct natriuretic action on the renal tubules, but this effect is overshadowed by its stimulatory action on the production of aldosterone by the adrenal cortex. This in turn promotes sodium reabsorption in exchange for potassium in the distal renal tubules. Renin itself cannot be measured easily, but the activity of the renin system can be gauged by the rate of production of angiotensin I from endogenous substrate during incubation of plasma in the presence of inhibitors of the converting enzyme and angiotensinas; this estimation is known as the plasma renin activity (PRA). A similar measurement is available known as plasma renin concentration, in which excess exogenous substrate is added eliminating the effect of substrate variation. In addition there are assays for determining the levels of circulating angiotensin II (Boyd, Landon, and Peart, 1967).

In children interpretation of plasma renin values has proved difficult because of the limited normal data available. Contributory factors in this dearth of paediatric information have been the large quantities of blood hitherto required for measurement of the various parameters of the renin-angiotensin system and the sampling difficulties encountered in young children. Several studies have suggested that in children peripheral venous plasma renin levels were higher than in adults (Godard et al., 1968; Amsterdam et al., 1969; Kotchen et al., 1972). Krause, Schillmoller, and Hayduk (1972) confirmed these findings utilizing a bioassay technique for plasma renin concentration and showed a significant negative correlation between renin concentration and body surface area. More recently, Dillon and Ryness (1974), using a semimicro radioimmunoassay have established normal ranges for peripheral plasma renin activity in infancy.
and childhood and observed a tenfold decline in PRA with age, falling from 1392 pg angiotensin I/ml per hr in infancy to 87 pg angiotensin I/ml per hr in adult life. Plasma aldosterone concentration similarly declines from 20·2 ng/100 ml to 11·7 ng/100 ml.

Sodium deprivation enhances renin secretion, and it is established that PRA should be interpreted in relation to sodium turnover (Laragh et al., 1972) or total exchangeable sodium (Davies et al., 1973). It is not obvious how to compare rates of sodium turnover in infants and adults but the decline of PRA with age cannot be wholly attributable to differences in sodium intake. A possible explanation is that the young infant is more dependent than the adult upon renin-aldosterone stimulated distal tubular sodium reabsorption for the maintenance of salt balance.

**Renal hypertension**

The conditions in which hypertension is associated with activation of the renin-angiotensin system include renal ischaemia due to renal artery stenosis, many varieties of parenchymal renal disease especially when hypertension is resistant to saline depletion (Weidmann et al., 1971; Schalekamp et al., 1973), rare renin-secreting tumours (Robertson et al., 1967), occasionally Wilms's tumours (Mitchell et al., 1970), and malignant hypertension in some individuals. Secondary hyperaldosteronism with hypokalaemia may occur in all these situations.

**Renovascular disease.** Renovascular hypertension can be defined as hypertension caused by a lesion of the renal artery or its branches which impairs blood flow to all or part of the kidney (Leumann et al., 1970). It is a curable disorder that has recently been recognized with increasing frequency. It is second only to coarctation of the aorta as a cause of surgically remediable hypertension in children (Fry et al., 1973). Renal artery stenosis with hypertension has been reported in neonates (Ljungqvist and Wallgren, 1962; Schmidt and Rambo, 1965; Angella et al., 1968; Formby and Emery, 1969). It is sometimes familial (Bergstein et al., 1971; Kaufman et al., 1972). Renal artery stenosis has been associated with idiopathic hypercalcaemia (Wiltse et al., 1966), Marfan's syndrome (Loughridge, 1959), the rubella syndrome (Menser et al., 1966), Takayashi disease (Fry et al., 1973), and neurofibromatosis (Halpern and Currarino, 1965; Bourke and Gatenby, 1971; Klecker and Roth, 1974). A variety of pathological lesions have been reported, including intimal hyperplasia, arteritis, and external compression. However, the commonest finding is fibromuscular dysplasia, particularly in older children (Fry et al., 1973). The disease may be bilateral, may involve segmental vessels (Kaufman et al., 1972; Fry et al., 1973), and occasionally the renal artery is replaced by many smaller vessels (Barratt, 1974).

Other causes of renovascular hypertension in children include renal artery aneurysm (Grossman and Babbit, 1967; Kaufman et al., 1972), arteriovenous fistula (Long, Javid, and Julian, 1964), intrarenal vascular anomalies (Leumann et al., 1970; Chrispin and Scatliff, 1973), and renal artery disruption after trauma (Fry et al., 1973).

The ischaemic kidney secretes excess renin but not all cases of renal artery stenosis have raised peripheral plasma renin activity (Brown et al., 1965) or angiotensin II concentration (Catt et al., 1971). On the other hand, Vaughan et al. (1973) found that if peripheral plasma renin levels were related to the urinary sodium excretion they were abnormally high in adults with renal artery stenosis. Dillon and Ryness (1974) have confirmed that most children with hypertension secondary to renal vascular anomalies or renal scarring have higher PRA than healthy children of equivalent age.

In terms of preoperative prediction of surgical cure it appears that it is not the peripheral plasma renin activity but the differential renal vein renin levels which are of greatest value (Michelakis et al., 1967; Stockigt et al., 1972). A renal vein renin ratio of greater than 1:1·5, especially if associated with evidence of suppression of renin release from the contralateral kidney, predicts a good response in terms of surgical treatment for renovascular disease and is useful in patients with other types of renal hypertension (Stockigt et al., 1972). However, renal vein renin determinations in children are even more prone to error than in adults. False negative ratios may occur for many reasons (Poutasse et al., 1973; Vaughan et al., 1973) and it must be remembered in this context that β-adrenergic blockers specifically depress renin release and should be withdrawn at least 3 days, but ideally 2 weeks, before this investigation is undertaken. Selective sampling from segmental veins draining underperfused areas of kidney may allow the identification of localized sources of renin secretion which may be overlooked by main renal-vein sampling (Schambelan et al., 1974). Successful revascularization of an ischaemic kidney in childhood offers a good prospect of permanent cure of hypertension (Kaufman et al., 1972; Fry et al., 1973).
Parenchymal renal disease. The relation between parenchymal renal disease and hypertension is very complex. Hypertension is regularly associated in children, as in adults, with advanced renal disease and uraemia. However, with some important exceptions, the renin angiotensin system cannot be implicated in this group of patients. In most of the patients the hypertension responds to salt depletion but in the remainder this fails to lower the blood pressure; these individuals have high circulating renin levels in their peripheral blood in relation to their exchangeable sodium (Schalekamp et al., 1973).

Lesser degrees of parenchymal renal damage may be implicated in the causation of renin-dependent hypertension in children. Included within this group are probably segmental renal hypoplasia (Ask-Upmark, 1929; Royer et al., 1971), the kidney after renal venous thrombosis (Perry and Taylor, 1940), and also children with localized pyelonephritic scarring and hypertension. In unilateral renal disease with negligible function on that side and with an apparently normal or hypertrophied contralateral kidney, nephrectomy may well be undertaken in the hope that the blood pressure will fall. In bilateral disease in which the disparity of function is less marked, then medical treatment is to be preferred (Barratt, 1974). Renal vein renin studies may well reveal clear lateralization of renin output in this type of case, but it is worth noting that with some pyelonephritic kidneys the hypertension sometimes subsides spontaneously without resort to operation (Barratt, 1974). No clear guidelines have been established in children for deciding which kidney should be removed and if in doubt, a conservative approach is preferable.

Renal tumours. It has been reported that Wilms's tumours are occasionally associated with hypertension (Koons and Ruch, 1940; Hughes, Rosenblum, and Horn, 1949) and that this has been associated with increased renin output from the offending kidney (Mitchell et al., 1970). Whether this is due to impairment of vascular supply to part of the kidney or to increased secretion of renin from the tumour itself is not entirely clear, but the findings suggest the latter (Lee, 1971). More recently it has become apparent that specific renin secreting tumours exist, known as haemangio-pericytomas. There have been 7 published reports (Robertson et al., 1967; Kihara et al., 1968; Eddy and Sanchez, 1971; Bonnin, Hodge, and Lumbers, 1972; Conn et al., 1972; Schambelan et al., 1973; Brown et al., 1973) and of these, 3 were in children, the youngest being a girl of 8 years. Characteristically, those affected have hypertension, evidence of secondary hyperaldosteronism, normal renal function, normal IVP, and a normal or only slightly abnormal renal arteriogram. Divided renal vein renin levels reveal clear lateralization to the side affected and nephrectomy results in cure. So far, there is no evidence of malignancy and no metastases have been reported.

In conclusion, it is becoming increasingly apparent that disturbances of the renin angiotensin system in childhood are of considerable importance. An understanding of the physiology and pathology is essential for the satisfactory diagnosis and treatment of many hypertensive disorders which affect children. With development of semimicro methods for measurement of plasma renin activity and more refined techniques of arteriography and reconstructive vascular surgery, there is every reason to expect improvement in the evaluation and treatment of these children.

This work was undertaken during tenure of the Alan Moncrieff Educational Research Fellowship supported by the Buttle Trust.

REFERENCES


Annals of the Academy of Medicine, Singapore, Vol. 11, No. 1, January-February 1982. Published by the Academy of Medicine, Singapore.


M. J. DILLON

The Hospital for Sick Children, Great Ormond Street, and Institute of Child Health, London WC1N 1EH.