Hypertension in the newborn baby

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Hypertension is rare in the newborn infant. It has a cumbersome definition and diagnosis, and screening is not justified using present definitions and technology. Thresholds for starting antihypertensive treatment in the first month of life are not clear, and the treatment is difficult, with idiosyncratic responses to drugs in neonates with varying renal and hepatic function.

Neonatologists diagnose and treat hypertension most days of their lives, but rarely diagnose hypertension in the newborn baby. Why? The definition of hypertension is based on centiles from population data. There are as many babies below the 5th centile as there are above the 95th centile, but only the former receive much attention. This paradox is heightened by recommendations that universal screening of blood pressure in neonates is not warranted. Therefore, blood pressure is measured only in the 7–8% of newborn babies admitted to neonatal units—usually screening for hypotension—and in another 1–2% of babies with a recognised cardiovascular or renal abnormality.

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DEFINITION

Hypertension in children and infants is diagnosed as a systolic and/or diastolic pressure $\geq 95$th centile for age and sex on three separate occasions, but in the newborn only systolic values are used. This epidemiological definition based on centiles has been criticised as having no biological meaning for children because it was introduced to assess adults and essential hypertension.

MEASUREMENT

The baby should be quiet, and not feeding; systolic pressure is 5 mm Hg lower in sleeping babies. In newborns, conventional sphygmomanometry is not recommended because the Korotkoff sounds cannot be heard reliably. Instead Doppler ultrasound, pulse oximetry, or oscillometry are used. The pulse oximetry technique uses the disappearance and reappearance of the plethysmographic waveform of a pulse oximeter as the cuff is inflated and deflated around systolic pressure. It has been used in infants in intensive care with distinct oximetry waveforms, but further work is needed to support its wider use. Oscillometric manometers are the most widely used instruments in clinical practice, but are less accurate than either Doppler or pulse oximetry techniques when compared with the gold standard of intra-arterial pressures. For non-invasive measurements, the cuff bladder should cover 80–100% of the upper arm circumference and its width should be 40% of that same circumference.

NORMAL DATA

The American Second Task Force on Blood Pressure Control in Children incorporated data from over 70 000 American and British children into its centiles. More recently, two large British studies produced reference ranges for preterm babies, recording mean blood pressure using indwelling arterial catheters, and systolic pressure by Doppler ultrasound. Neither reported the 95th centiles, the closest being the 97th centiles shown graphically in the latter, which also included data on term babies. Table 1 summarises these findings.

SCREENING AND TRACKING

The original criteria of Wilson and Jungner for disease screening programmes have been enunciated by the National Screening Committee. In brief, the disease itself should be an important problem, with a known natural history that progresses from a latent to a declared phase with a detectable risk factor or disease marker in the former. There should be an effective treatment. The test should be validated, simple, safe, and precise with an agreed cut off value. It should be acceptable to the population being screened and be economically balanced. Hypertension in the newborn does not fulfil several criteria. The difficulty and time costs of three separate accurate measurements do not equate to "a validated simple, safe, and precise screening test". Also, neonatal hypertension does not "progress from a latent to a declared disease". Tracking is the concept that a child will grow along (track) his blood pressure centile into adulthood—that is, a child with a blood pressure on the higher centiles will develop into a similarly placed adult. This correlation increases with age, but in the Brompton study "at ages 4 days, 6 weeks and 6 months blood pressures were in effect not correlated with blood pressures later in life".

INCIDENCE AND AETIOLOGY

The reported incidence of hypertension in neonates varies from 0.2% in the oft quoted data from Ingelfinger for healthy newborns, up to 2.6% in babies after neonatal intensive care, and 40% in patients with chronic lung disease. However, many reports used non-standard definitions, had a biased selection of babies, and an incomplete...
follow up. One group concluded that “at most only 1% of children have blood pressures consistently and appreciably above the 95th centile for single measurements”.

Table 2 shows data from some papers on hypertension in babies on or recently discharged from neonatal units. Three American series reported incidences of 0.7%, 0.81%, and 2.0% of hypertension on their neonatal intensive care units, but different thresholds for diagnosing hypertension were used. Some 80% of their hypertensive babies had had an umbilical artery catheter in situ before the hypertension. This was a risk factor when hypertension was diagnosed in the first 2 weeks of life, but not when it was diagnosed at follow up (table 2). Improved catheter design and the use of heparin have reduced but not eradicated this problem.

Babies with chronic lung disease can become hypertensive. Dexamethasone treatment of chronic lung disease raises systolic blood pressure by a median of 27 mm Hg and this usually resolves two weeks after treatment is stopped. Pain, patent ductus arteriosus, extracorporeal membrane oxygenation (ECMO), other renal causes, endocrinological disorders, maternal drug abuse in pregnancy, tumours, and neurological problems should all be considered.

INVESTIGATION

Recognition of specific conditions such as those listed above may limit the need for further tests. The baby should be re-examined with particular attention to systolic pressures in the upper and lower limbs, murmurs, bruises, and abdominal masses. Signs of end organ damage should be sought; retinopathy, encephalopathy, left ventricular hypertrophy, haematuria, and proteinuria must all be considered.

Basic plasma and urine biochemistry should be checked. All hypertensive babies should have an ultrasound scan of the renal tract and suprarenal region, and the abdomen should be radiographed by an experienced paediatric radiologist. Infants who have had an indwelling umbilical artery catheter should have their aortas and renal arteries imaged, at the very least by ultrasonography, to look for thrombi. These and renal vein thromboses, congenital abnormalities of the renal tract, obstructive uropathies, parenchymal renal disease, and tumours may be seen. Doppler studies of the renal arteries may reveal stenosis, but branch artery stenosis can be missed. The brain should be scanned for intracranial haemorrhage and cerebral oedema. Renograms, plasma renin activity, aldosterone, cortisol, and urinary steroid profile, urinary catecholamines, or thyroid function may be appropriate for selected patients.

OUTCOMES

Neonatal hypertension improves during infancy but the nature of this trend is obscured by variations between studies (table 2). Sheftel et al diagnosed hypertension in seven (9%) of 79 preterm babies who had been in intensive care in the first three months after discharge. Follow up was for only 21 weeks, and by then three babies had had surgical treatment, and three of the four “idiopathic” hypertensive babies were
still receiving antihypertensive drugs. Buchi and Siegler reported 53 babies who were hypertensive on a neonatal unit by their own lower thresholds of systolic pressures (> 90 mm Hg for term infants and > 80 mm Hg for preterm infants). By 12 months, 81% were normotensive, but two had died from the complications of hypertension, and in four it persisted. Friedman and Hustead followed 17 (2.6%) of 654 preterm babies who developed hypertension after discharge from a neonatal intensive care unit. Sixteen were treated, four surgically. Propranolol was the only drug used in all but one, who also had chlorthalidzide. All babies were off therapy by 24 months. The earlier group of nine infants of Adelman et al developed systolic pressures ranging from 115 to 280 mm Hg on a neonatal unit. Eight had renal artery thrombi in association with umbilical artery catheters. One was treated with chlorthalidone alone, seven with hydralazine and methyldopa, and one with propranolol alone. Treatment was discontinued in all by seven months, and they remained normotensive. These generally favourable outcomes are reassuring, but there are reports of death, heart failure, encephalopathy, and retinal changes in some hypertensive babies. Neonatal hypertension is not entirely benign.

TREATMENT
First, do no harm!

The received wisdom is that an asymptomatic neonate with a systolic pressure consistently between the 95th and 99th centiles and with no end organ involvement should be observed but not treated, in the expectation that the hypertension will settle. The dilemma is between “unnecessary” medication and its potential side effects for a condition that is likely to resolve, and a fear of sudden life threatening end organ damage secondary to untreated hypertension. The general paediatrician may therefore wish to consult a paediatric nephrologist with greater experience of this problem in young infants, particularly as many cases will be secondary to renal disease. Feld and Wax advocate treatment if the systolic pressure is above the 99th centile or if there is end organ involvement with a systolic pressure above the 95th centile. The Task Force’s 99th centile was 110 mm Hg on day 7 of life, but the 97th centile in a recent study was 111 mm Hg on day 10. There is little if any evidence for a precise, single starting point for treatment. A blood pressure more than 30% above that expected for age can be considered as a hypertensive emergency. Empirically therefore one might treat a week old baby with a systolic pressure consistently above 110–115 mm Hg using a Doppler technique, and regard a systolic pressure of > 130 mm Hg at that age as a hypertensive emergency.

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Treatment should proceed in a stepwise fashion. First reduce or withdraw treatment that may raise blood pressure. Salt or fluid overload must be corrected. If investigations reveal a surgically remediable cause, the timing of surgery depends not just on the nature of the lesion but also on the severity of the hypertension, the benefits and risks of medical treatment, and the age and weight of the baby. If medical treatment is needed, the choice is between five groups of drugs: diuretics, β-adrenergic blockers, calcium channel blockers, direct peripheral vasodilators, or angiotensin converting enzyme inhibitors. Their use has to be tailored to the cause, to any impending hypertensive crisis, and to the routes available for administration in a particular baby. Many of the drugs are not licensed for children, have different pharmacokinetics in neonates, and experience with them in this age group is limited. All drugs should be started at their lowest doses, as found in standard texts.

Diuretics increase salt and water excretion, leading to decreased extracellular and plasma volumes. Although compensatory mechanisms come in to maintain sodium homoeostasis and plasma volume may return towards normal, there is a sustained reduction in volume and a modest fall in blood pressure. Unless a specific problem with high volume hypertension is recognised, their use in neonatal hypertension is limited. Indeed, if volume overload is not present, they may contribute to a hypotensive crisis if used with other antihypertensive drugs.

Propranolol has the advantage of being the most extensively used β-blocker in neonates with hypertension, and has a low incidence of side effects. Labeltol blocks both β and α receptors, although it is eight times more potent against the former. Pure α-adrenergic antagonists are rarely used in neonatal hypertension.

Nifedipine is the type II calcium channel blocker that has been used most in babies. It has a vasodilator action that low- ers peripheral vascular resistance. Hydralazine and minoxidil act directly on the vascular smooth muscle to reduce peripheral vascular resistance. Hydralazine is the first choice clinically as the more potent minoxidil has a number of adverse effects, and is reserved for those with refractory hypertension. All the vasodilators may initially cause an increase in heart rate and cardiac output and flushing. Captopril, an angiotensin converting enzyme inhibitor, appears to be more potent and longer acting in the neonate than older children. Doses as low as 10 µg/kg (a tenth of the normal low starting dose) may be effective. Renal vascular resistance is high in neonates, hence the effects of angiotensin converting enzyme inhibitors are more pronounced. If renal vascular disease is suspected, its use should be avoided until it is clear that the renal vasculature is normal. Neurological complications and oliguria have been reported after administration of captopril to neonates and it should be used cautiously only after an initial test dose.

In a hypertensive crisis, the intravenous drug of choice is labetalol or hydralazine. These can be titrated against the blood pressure, which may begin to fall within an hour of administration. Some advocate diazoxide but the concern in neonates, and experience with them in this age group is limited. All drugs should be started at their lowest doses, as found in standard texts.

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SUMMARY
Hypertension is rare in the newborn infant. It has a cumbersome definition and diagnosis. Screening is not justified using present definitions and technology. There has been little work in the 1990s on hypertension in extremely preterm babies. Thresholds for starting antihypertensive drug treatment in the first month of life are unclear and debatable. Treatment is difficult, with idiosyncratic responses to drugs in neonates who have varying renal and hepatic function. There is a need to keep registers of babies who are hypertensive in order to audit the outcome of management with or without antihypertensive drugs, and to establish appropriate thresholds for starting treatment.

REFERENCES
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