Lung transplantation: technical problems

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The biological problems involved in successful organ transplantation are well known and have been reviewed by Gowans (1962) and by Howard and Michie (1963). Besides these there are many technical difficulties.

These may be studied experimentally by lung excision and re-implantation in the same animal, or by lung transplantation from unrelated or related animals. Further, the tolerance time of lungs to anoxia can be evaluated.

LUNG EXCISION AND RE-IMPLANTATION

In 1951, Juvenelle, Citret, Wiles, and Stewart reported a six-month survivor from a series of dogs subjected to pneumonectomy and re-implantation of the vessels and bronchi of the right lung. In 1952, Neptune, Redondo, and Bailey had a successful re-implant in dogs one year after operation. In 1959, Huggins reported successful excision and re-implantation of the left lower lobe of the dog lung. In 1963, Alican and Hardy reported 33 long-term survivors in dogs in which either the right or the left lung had been re-implanted.

In studying these survivors further by contralateral pneumonectomy, pneumonectomy and re-implantation, pulmonary artery ligation or multiple lobectomies, they drew the following conclusions:

1. Pulmonary oedema may frequently develop when the blood-flow through the re-implanted lung is suddenly increased.

2. The altered pattern of respiration due to the absence of reflexes originating in the lungs may not be compatible with long-term survival, or at least the animal without these reflexes is at a definite disadvantage as far as ventilatory adjustments are concerned.

3. In future lung homotransplantations, the preservation of an appropriate amount of the recipient's own lung tissue to prevent alteration in the normal respiratory pattern and to decrease the chances of pulmonary oedema would be of serious concern.

EXCISION AND RE-IMPLANTATION OF LUNGS IN SHEEP

Since 1957 the technical steps that have to be taken in transplanting lungs have been studied in the Department of Surgery, University of Otago. The purpose of this paper is to describe the techniques used and the problems encountered.

Initially, because of the homograft reaction, the stages in a suitable technique were established by completely excising and re-implanting a lung of the same animal, checking the efficacy of the procedure by differential lung function studies of oxygen uptake before operation, immediately afterwards, and again after two months. It was felt that the post-operative ability of a lung to absorb oxygen as determined by differential bronchospirometry gave the best index of the success of the experiment. Histological studies were confirmatory. Further, such a procedure could give a completely denervated lung, which is ideal for physiological study of the nature of respiratory reflexes.

The sheep was chosen as the experimental animal, and the left lung was used throughout. The right lung was deemed unsuitable because of its tracheal bronchus to the cephalic lobe.

LUNG FUNCTION: DIFFERENTIAL BRONCHOSPIROMETRY

For experimental ease, readily available standard equipment was used. Though the trachea of a sheep is too long to use a normal Carlens endobronchial tube inserted through the mouth, pre- and post-operative bronchospirometry as a test of lung function by oxygen uptake can be satisfactorily performed through a temporary tracheostomy (Borrie and Montgomerie, 1958a and b).

Animals weighing from 40 to 45 kg. were used. After premedication with morphine (15 mg.) and atropine (1·2 mg.), general anaesthesia was induced with pentobarbitone (64 mg./8 kg. body weight intravenously), supplemented as required with open ether. The trachea was exposed below the thyroid gland and incised transversely. A no. 37 or 39 Carlens tube was inserted, and, with the animal still lightly anaesthetized and breathing regularly, pul-
monary ventilation and oxygen uptake were recorded with a Palmer twin-recording bronchospirometer.  

Once a satisfactory record had been obtained, the Carlens tube was removed, a size 8 cuffed Magill endotracheal tube was inserted through the tracheostomy, and anaesthesia was continued by attaching the tube to a Palmer anaesthetic pump which had been set to the appropriate tidal volume as determined from a weight–tidal volume chart (Borrie and Mitchell, 1960). 

After completing the experimental procedure, the chest wall was closed over water-seal drainage, and post-operative differential bronchospirometry was performed by removing the Magill tube and re-inserting the Carlens tube. Finally, the tracheostome was closed with interrupted sutures placed through the peritracheal fascia. It had previously been shown that simple transverse tracheal wounds in the sheep heal rapidly without stricture formation (Borrie, 1957). Each animal also received penicillin, 1 million units, and streptomycin, 1 g. daily for seven days after operation.

Two months after operation lung function was reassessed by repeating the procedure. Thereafter the animal was killed, and symmetrical specimens from the two lungs were taken to determine the normal sheep lung histology.

TECHNIQUE OF LEFT LUNG EXCISION AND RE-IMPLANTATION

After several pilot experiments total lung excision and re-implantation was performed (Borrie and Montogomerie, 1958a and b).

Excision After differential bronchospirometry a standard left thoracotomy was performed through the fifth intercostal space (Fig. 1). The pulmonary liga-

![FIG. 1. View of the medial wall of the left pleural cavity of a sheep via a left fifth intercostal thoracotomy.](http://thorax.bmj.com/content/15/3/384.sa)

![FIG. 2. View after incising the mediastinal pleura and ligating the left azygos vein.](http://thorax.bmj.com/content/15/3/384.sa)

![FIG. 3. Technique of lung excision.](http://thorax.bmj.com/content/15/3/384.sa)
The pericardial flaps. The hemiazygos vein, a constant feature of the sheep, was next ligated and divided as it entered the coronary sinus (Fig. 2).

The left main bronchus was isolated from the level of the carina to its upper lobar branch, a 2-cm. length, carefully clamped at the carina, and divided (Fig. 3a).

The left pulmonary artery was dissected to its origin, doubly clamped with Potts spoon-shaped multi-toothed arterial clamps, and divided (Fig. 3b).

The upper and lower left pulmonary veins were finally isolated, clamped with a toothed bronchus clamp, and divided, thus excising the lung (Fig. 3c).

After excision of the lung these venous openings into the heart were closed in two layers with continuous silk sutures. The artery of the excised lung was immediately perfused with 6% dextran in 0-9% saline at 4° C. and 15 mm. Hg pressure.

RE-IMPLANTATION It was initially assumed to be technically easier and more satisfactory to anastomose the common left pulmonary venous stump to a fresh opening in the left auricle than to rejoin it to the pulmonary venous stumps. A suitable length of auricle was therefore lifted in a Brock auricular clamp and incised, and a one-layer vascular anastomosis was made with Blalock evertting sutures between the pulmonary veins and the heart. Later experience, however, with re-implantation of the lungs showed it was difficult in sheep to get an adequate cuff of atrium containing both left pulmonary veins for this re-implantation experiment, and that it was desirable to resuture the veins to their original site.

The pulmonary artery was similarly resutured (Fig. 4a), and the clamps on the artery and vein were released. The average time taken to complete this vascular stage was one and a quarter hours.

Thereafter, suture of the left main bronchus completed the re-implantation (Fig. 4b). Additional air-seal was secured by a free fascial graft collar around the bronchial suture line, taken from the serratus anterior muscle. The chest wall was closed with water-seal drainage, and further spirometry was performed to assess the oxygen uptake per lung.

Results None of the 10 sheep in the original series survived more than six days; deaths within 24 hours were due to tension pneumothorax (1), operative haemorrhage (2), and bronchial obstruction (3), and later deaths were due to a slowly developing thrombosis within the pulmonary vascular bed, producing pulmonary infarction.

Of five animals investigated by immediate post-operative differential bronchospirometry, two showed ventilatory and oxygen absorbing function similar to the values found pre-operatively (Fig. 5a and b), and one showed a latent function made apparent when the right lung was occluded (Fig. 5c and d).

Comment These experiments showed that it was possible to restore ventilatory and oxygen absorbing function, at least temporarily, after excising and re-implanting a lung in sheep, and raised the hope that longer survivals could successfully be achieved. Death was most commonly due to pulmonary infarction. As this could have arisen in the pulmonary or bronchial arteries, or the veins, its basic cause and method of prevention were further investigated by studying separately the effects of dividing and resuturing the four major components of the lung hilum. Each of these problems was checked in sets of six experiments controlled by lung function studies before, immediately after, and up to three months after operation.

1. BRONCHIAL ARTERY DIVISION AND LUNG FUNCTION

In the earlier 'total' experiment of lung excision and re-implantation the minute left bronchial artery was merely ligated. The effect of this ligation on subsequent lung function was therefore first investigated.

Technique After differential bronchospirometry the hilum of the left lung was dissected, the left main bronchus isolated, the bronchial artery divided and ligated, and the thoracotomy closed. Differential bronchospirometry was then repeated, and again two months later, after which the sheep were killed and the lungs were examined histologically (Fig. 6).
FIG. 5. Differential spirograms of sheep showing oxygen uptake of the left lung (a) before and (b) immediately after total excision and re-implantation. (These spirograms are read from right to left.) Similar spirograms taken (c) pre-operatively and (d) post-operatively show a latent function in the left lung made more obvious when the right lung is temporarily occluded for minute intervals.
FIG. 6. Histological pattern of (above) the right and (below) the left lung two months after division of the left bronchial artery.
Results All six animals survived operation. Preoperative spirometry established the normal oxygen uptake of each lung. Immediately after operation the total oxygen uptake was reduced, but the left lung/right lung oxygen absorption ratio remained unchanged in all but one animal. Two months later there was still the same proportionate oxygen uptake by the left lung, and the animals remained healthy and thriving (Fig. 7).

It is concluded that in the normal sheep the rate of oxygen uptake is greater in the right than in the left lung, and that division and ligation of the bronchial artery has no measurable effect on the proportionate oxygen uptake of the left lung either immediately after or within two months of operation (Borrie, Campbell, and Fulton, 1958).

2. PULMONARY ARTERY DIVISION, RESUTURE, AND LUNG FUNCTION Normal pulmonary vascular pressures in the sheep were measured. They were: pulmonary artery, 20/14 mm. Hg; pulmonary vein, 5/2-25 mm. Hg; and left atrium, 7-5 mm. Hg. When both pulmonary veins were clamped, the pressure on the pulmonary side rose and oscillated between 17 and 15 mm. Hg.

Technique After routine anaesthesia, differential bronchospirometry, and left thoracotomy, the left pulmonary artery was isolated, doubly clamped with Potts' arterial clamps, divided, and then resutured using the Blalock evertting technique with 4-0 silk sutures. Anticoagulants were not used. Post-operative bronchospirometry was performed, and the wounds were closed. Three months later, after further differential bronchospirometry, the lungs and pulmonary arteries were examined.

Results All eight animals survived operation, but two died in the early post-operative period, from sputum retention, and from empyema.

Immediate and late post-operative bronchospirometry showed that the proportionate oxygen uptake of the left lung was unaltered by the operation (Fig. 8). Necropsy confirmed patency of the left pulmonary artery in all cases and revealed a soundly healed anastomosis with no evidence of intravascular clotting or obstruction (Fig. 9). The histological picture was normal in the test lung.

FIG. 7. Differential spirograms before, immediately after, and two months after operation, showing satisfactory oxygen uptake in the left lung after division of the bronchial artery.

FIG. 8. Differential bronchospirometry where the left pulmonary artery was divided and rejoined: tested before, immediately after, and two months after operation.

FIG. 9. The left pulmonary artery showing the site of division and resuture and no occlusion or narrowing of the lumen two months after operation.
It is concluded that division and resuture of the left pulmonary artery is regularly feasible without interfering with the ability of the left lung to absorb oxygen. These results further suggested that the initial problem of intravascular clotting after lung excision and re-implantation did not lie on the arterial side (Borrie and Fulton, 1958).

3. LEFT MAIN BRONCHUS DIVISION, RESUTURE, AND LUNG FUNCTION

Technique Once again, after anaesthesia, differential bronchospirometry, and left thoracotomy, both the right and left main bronchi were isolated in the mediastinum. The left main bronchus was dissected free from the carina down to its upper lobe bronchus. It was lightly clamped on the proximal side only at carinal level with a Satinsky clamp, and divided in the middle of the exposed segment. Resuture was performed with a continuous 0000 silk suture, using the Blalock technique.

An interrupted locking stitch was also used after every three bites of the continuous suture. After the resuture had been completed, the clamp on the bronchus was released, and the lung was inflated. A fascial collar 2 cm wide was again cut from the serratus anterior muscle, threaded round the line of anastomosis, and sutured into place. This achieved air-tight bronchial closure. The chest was closed by standard technique with tube drainage. Post-operative bronchoscopy and differential spirometry were next performed, and the tracheostome was closed.

Two months later, under general anaesthesia, bronchoscopy was performed, followed by differential bronchospirometry. The lungs and bronchial suture line were later examined.

Results The experiment was performed on six sheep without death. In the first animal, in which the technique was being established, there was a residual left bronchial stricture, but some oxygen uptake of the left lung. The remaining five animals two months after operation showed normal oxygen uptake in the test lung (Fig. 10). Necropsy showed a normal bronchial lumen with no suggestion of narrowing at the line of bronchial resuture (Fig. 11). The histology of the five lungs was also normal (Fig. 12).

It is concluded that this technique gives a satisfactory bronchial anastomosis after bronchial division, without air leak, stricture formation or adverse effect on lung function up to two months after operation (Borrie and Foster, 1959a).

4. DIVISION AND RESUTURE OF PULMONARY VEINS

In the early reported successes in dogs of pulmonary re-implantation and transplantation, a left atrial cuff containing the orifices of the two pulmonary veins had usually been re-anastomosed to a new opening fashioned in the left atrial appendix.

In sheep, however, with a small, squat left atrium, a mechanical problem arose. For, even when using anticoagulants, the short length of
FIG. 12. Histological pattern of (above) the right, and (below) the left lung two months after division and resuture of the left main bronchus.
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Pulmonary veins available for anastomosis to the atrial appendix, and consequent stitching, caused collapse and mechanical occlusion of the venous lumen, leading to the rapid onset of venous thrombosis with pulmonary infarction (Borrie and Foster, 1959b).

Further, whereas venous autografts regularly remained patent, intravenous thrombosis regularly occurred after the experimental use of freeze-dried venous homografts.

In the further experiments, therefore, in re-implantation of lungs in sheep, the pulmonary veins were re-implanted into their normal position. The procedure was greatly helped with a Sellors atrial clamp.

**Technique** After differential spirometry and left thoracotomy the lung hilum was dissected. The left bronchial artery was ligated, the pericardium incised, the pulmonary artery encircled with a tape, and both left pulmonary veins dissected. Twelve thousand i.u. of heparin were given, and then the circulation to the left lung was interrupted by tightening the arterial tape and clamping both veins well on to the left atrium with a Sellors atrial clamp. Both pulmonary veins were divided and in turn resutured, using everting sutures of 0000 silk with every second stitch locked.

Pulmonary circulation was re-established, protamine sulphate was injected to neutralize the heparin, the chest wall was closed, and post-operative bronchospirometry was performed.

**Results** Of four sheep in which the operation was performed, three survived. In the surviving sheep, post-operative spirometry showed normal oxygen uptake in the left lung before, after, and two months after division and resuture of the left pulmonary veins.

**FIG. 13.** Differential spirogram showing normal oxygen uptake in the left lung before, after, and two months after division and resuture of the left pulmonary veins.

**FIG. 14.** Macroscopic appearance of two left lungs with normally patent pulmonary veins two months after division and resuture.
FIG. 15. *The normal histological lung pattern of one of these sheep two months after operation: (above) the right lung; (below) the left lung.*
completed, three had normal oxygen uptake three months after operation (Fig. 13). Macro-
scopically the anastomosis was well healed with-
out any narrowing (Fig. 14). Histologically these
lungs were normal (Fig. 15). The fourth animal
survived, but at the anastomosis, which was nearer
the lung, there was a stricture, and the lung
contained a large abscess.

Conclusion Successful pulmonary venous anas-
Mosis, when re-implanting a sheep lung, though
difficult, is feasible. The lungs absorb oxygen
normally and are histologically normal three
months after operation. For success the left
pulmonary veins should be divided and resutured
as close as possible to the left atrium at the normal
site of entry of these veins. This in turn requires
a satisfactory atrial clamp and a careful locking
suture technique (Borrie and Prapaiwongs, 1960).

TOLERANCE TIME OF SHEEP LUNGS TO
ANOXIA

How long can a lung tolerate total anoxia yet
remain viable and function normally? This is a
vital question in any lung transplantation experi-
ment or clinical procedure, as an index of the time
available for re-establishing pulmonary blood flow.

METHOD

After differential bronchospirometry and left thora-
cotomy under normothermia, the pulmonary ligament
was divided, the animal heparinized, the bronchial
artery ligated, the left main bronchus clamped, and
the pulmonary artery and both veins were snared and
then firmly occluded.

In successive experiments the anoxic period was
increased by quarter-hour intervals from one hour up
to four hours.

At the end of each test period the bronchus and
vessels were released, the circulation was re-
established, the left lung inflated, and the heparin
neutralized with protamine sulphate. The chest wall
was closed, and differential spirometry was again per-
formed. In surviving animals, spirometry was repeated
two months later, after which the animals were killed
and the lungs were examined histologically.

RESULT

Eighteen experiments were performed. Nine of
the animals were subjected to periods of up to
two and a half hours of hilar occlusion, and all
survived with lungs functioning normally both
immediately and two months after operation.

Of the nine animals whose lungs were anoxic
for three hours or more, there were three survivors
after three, three and a half, and four hours of
anoxia respectively. The four-hour survivor
developed transient left pulmonary oedema, which
was successfully treated by endobronchial aspira-
tion for one hour and tracheostomy for 24 hours.
All three had normal oxygen uptake two months,
and histologically normal lungs three months after
operation.

CONCLUSION

It is concluded that the left lung of sheep can
regularly withstand two and a half hours of
anoxia, and that there is no sharp end-point to
tolerance of anoxia between three and four hours
of circulatory arrest in the test lung. After four
hours of anoxia with normothermia pulmonary
oedema will regularly occur (Borrie, 1962).

One has therefore at least two hours in hand
for transplanting lung, i.e., from the time of arrest
of circulation through the lung for transplantation
until it is once more receiving blood in its new
situation within the host.

LUNG HOMOTRANSPLANTATION

In 1947, the Russian surgeon Demikhov first
transplanted the lower lobe of the right lung of
one dog into the lung of another dog, with survival
of the recipient for four days. This work only
became widely known in 1962 (Demikhov, 1962).
That animals can survive after lung transplantation
was shown by Hardin and Kittle (1954) who, after
transplanting the left lung, immediately performed
a right pneumonectomy. Two animals survived for
six and nine days, proving unequivocally the
functional ability of the homologous lung during
these periods.

Experimenter in the 1950s showed that homog-
rafts were always rejected and that the recipients
usually died within 10 days of operation from
pneumonia, the end result of the homograft
reaction (Davis, O'Conor, Coloviras, and Strawn,
1952; Neptune et al., 1952).

Attempts to modify this reaction led Neptune,
Weller, and Bailey (1953) to give corticotrophin
to four dogs with lung homografts, thereby
increasing the average survival time to 25 days.
One dog survived 42 days. Hardin and Kittle
(1954) had survivals from 12 to 18 days after
cortisone. They also reported survivals up to 36
days when litter mates were used.

Again, the discovery that antimetabolites and
alkylating agents could suppress the immune
response in skin and kidney homografts led to
their use in experimental lung transplants.
Blumenstock, Collins, Hechtman, Thomas, and Ferrebee (1962) showed that methotrexate could prolong survival in dogs with a lung homograft up to 410 days, though serial chest films in the long-term survivors showed progressive opacification of the transplanted lung. Hardy, Webb, Dalton, and Walker (1963) found azothioprine (Imuran) to be the most active immuno-suppressant, though their average survival time was only 14 days.

MacPhee and Wright (1964) reported successful lung homografts in four dogs without operative or early post-operative mortality. Cyclophosphamide in combination with other drugs suppressed the immune response for periods up to 49 days. In three dogs, intravascular thrombosis at the atrial suture line was the cause of death.

De Bono and Brock (1964) also briefly reported the survival of dogs up to eight months after lung transplantation, and used methotrexate to suppress the immune response.

Further, the first case report of lung transplantation in man has now been published and, though the patient died 18 days after operation in renal failure, the transplanted lung was functioning well, with no evidence at necropsy of rejection of the graft (Hardy, Eraslum, and Dalton, 1963).

We studied the problems of lung homografts in twin lambs in 1962 and in twin calves in 1963.

HOMOGRAFTS IN TWIN SHEEP

These lambs, born in the New Zealand spring of 1961 (September), were marked at birth and reared to 100 lb. hoggets. Twin sheep, however, are rarely if ever monozygotic. The lung transplantation was completed in four hoggets and served to establish the technique in subsequent lung homografting in twin calves.

The findings showed initial oxygen uptake and good initial air entry by auscultation, the animal at first being fit and feeding normally. Within six days, however, the animals sickened, developed a left pleural effusion, and died on the seventh to ninth post-operative day from bilateral bronchopneumonia. Sectioning of the lungs showed a red, fleshy 'homograft' lung, intact lines of anastomosis, thrombosis in the pulmonary veins arising on the atrial suture line, and pneumonia in the non-grafted lung.

HOMOGRAFTS IN 'IDENTICAL' TWIN CALVES

Because of the failure with twin sheep, the experiments in 1963 were taken a stage further and performed on animals known to be monozygotic identical twins, namely calves.

These animals were obtained by air transport from the New Zealand Department of Agriculture Ruakura Animal Research Station, Hamilton, some 600 miles to the north. This station is in the centre of a huge dairy industry based on a cow population of over two millions, so that identical sets of twin calves are common, and have been collected and closely studied at Ruakura for over 18 years. This station has a celebrated herd of identical twin cows covering many research projects, so that the station field officers are well versed in selecting them. The criteria of selection are set out in the station report Notes on the Identification of Identical Twin Calves and in Hancock's papers (1952).

These include careful appraisal of the physical features including coat colour, the eye-lashes, ear fringes, and tail switch, the head shape including the lower jaw and muzzle, the body conformation, and pigmentation pattern. Particular attention is paid to comparing the pigmentation on the ears, nose, palate, tongue, lower jaw, lower lip, anovulval area, and udder. In these experiments we did not do confirmatory skin graft tests. Throughout we had the utmost co-operation from Mr. A. H. Carter, Senior Animal Geneticist, Ruakura Animal Station.

It was appreciated that methotrexate or Imuran might favourably affect the outcome of the experiments, but the makers of these drugs when approached were unable to deliver supplies because, they said, all that was available was required for kidney transplantation in man.

OPERATIVE TECHNIQUE

Stage 1: The donor With the animal held on its haunches, the first twin or donor was anaesthetized with pentobarbitone sodium on the basis of 64 mg. to 8 kg. body weight, and 40 mg. of suxamethonium dichloride was injected into the external jugular vein. The animal was then placed on the operating table, and the trachea was intubated with a large size Magill endotracheal tube.

With strict aseptic technique a left thoracotomy was performed and the rib retractor was inserted. A no. 11-gauge French needle connected to a blood-taking set was next inserted through the pericardium into the left ventricle, and two units of blood were collected in standard citrate medium. The animal was then heparinized, the trachea was cross-clamped to avoid soiling, and the heart and lungs were immediately removed.

The left or test lung was next dissected, defining in turn all hilar structures. The left main bronchus was
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lightly clamped and divided at carinal level. The left pulmonary artery was dissected and divided at its point of origin from the main pulmonary artery. Care was taken with the right pulmonary veins which were dissected, ligated near their point of entry into the left atrium, and divided. Thereafter the left lung was again lifted up, and, taking a generous cuff of atrium that included the entry of both the right and left pulmonary veins, the heart was removed.

The left lung thus had a long cuff of left atrium, left pulmonary artery, and left main bronchus. These were trimmed, and the lung, whose vessels contained heparinized blood, was temporarily placed in a sterile dish in a refrigerator at 0°C.

Stage II: The recipient The second twin was similarly anaesthetized, and a left thoracotomy was performed. The left main bronchus was lightly clamped at carinal level and divided just proximal to its first branch. Similarly, the left pulmonary artery was clamped with a Satinsky clamp and divided. As the pulmonary veins were not required for the re-implantation, their cardiac ends were ligated and divided. The pneumonectomy was thus completed in about 40 minutes from the start of stage I.

In these experiments, the two stages followed each other. Though highly desirable, in our experimental unit it was not possible to perform the two stages simultaneously. This, in turn, may have adversely affected our final results.

Stage III: The transplantation A generous left atrial cuff was taken in a Sellors atrial clamp, and an elliptical segment of atrial wall was removed.

Re-implantation was begun on the venous side by anastomosing the donor atrial cuff to the elliptical atrial incision of the recipient. Blalock continuous evertting sutures (0000 black silk) were used throughout with the interrupted locking mattress sutures already described.

The pulmonary artery was next anastomosed by the established technique. The Sellors clamp was then removed from the left atrium, the Satinsky clamp from the left pulmonary artery, and circulation through the transplanted lung was thus re-established. Because of the heparin given immediately before the donor animal was killed, there was little chance of clotting occurring in the capillary system of the donated lung. Immediately the clamps had been released the lung flushed red as blood again entered the pulmonary blood vessels.

The time from taking the donated lung until circulation was re-established through it in the recipient animal was approximately one and a half hours, which is well within the tolerance time of lungs to anoxia.

The chest wall was closed over water-seal drainage which was removed when consciousness returned. In addition, two polythene 'Readivac' catheters were inserted through the chest wall for subsequent aspiration of post-operative pleural effusion with Readivac flasks, sutured to the back, and firmly clamped.

If blood replacement was required this was now given into the external jugular vein using the units of blood previously collected from the twin.

POST-OPERATIVE CARE. The animals received careful nursing, the team visiting at least hourly during the day and three-hourly at night. On each occasion the Readivac flask was released and the pleural cavity thus aspirated.

On the first day, up to 100 ml of clear fluid could be aspirated at a time, but this quickly reduced to a few millilitres per day. The Readivac catheters were removed after three to four days.

FUNCTIONING OF THE LUNG Because of the previous long experience with sheep lung implantation, the finding that oxygen uptake was present in the lung at the conclusion of the operation, the knowledge that occasionally animals had succumbed during differential spirometry at the end of the procedure, and the wish to have survivors that could be tested for differential spirometry 20 or more days after operation, in this set of experiments differential spirometry was deliberately omitted at the conclusion of the operation.

RESULTS The operation was completed in three calves.
One died from bronchial stenosis seven days after operation.
Two returned to pasture at the animal farm some two weeks after operation. At that time there was good air entry in the transplanted lung. The animals thrived, gambolled freely in the fields at the farm, and ate greedily.
They were rechecked 55 and 48 days after operation by chest films, bronchoscopy, and necropsy.
Chest films at this stage showed opacification of the grafted lung.
Bronchoscopy showed an occluded left main bronchus half an inch beyond the carina.
Necropsy revealed that in each animal the transplanted lung had become completely rejected and that this test lung, now necrotic, was enclosed in a thick fibrinous pleural cast.

SUMMARY AND CONCLUSIONS

The technical problems of left lung excision and re-implantation in sheep, followed by left lung transplantation in twin calves have been studied over the past seven years.
A method for performing differential bronchospirometry with standard Carlens tubes and using temporary tracheostomy is described.
Early experiments on left lung excision and re-implantation in sheep showed that, although the re-implanted lung functioned for a time, acute infarction of the pulmonary vessels occurred. None survived more than six days.

Further analysis by sets of experiments on the left lung showed that (1) ligation of the bronchial artery in the lung did not affect the proportionate oxygen uptake either immediately or within two months of operation; (2) division and resuture of the left pulmonary artery is repetitively feasible without affecting oxygen uptake in the left lung; (3) the left main bronchus can be repetitively divided and rejoined without stenosis using the Blalock evertting continuous suture technique combined with interrupted locking sutures; (4) successful pulmonary venous anastomosis is feasible and the lungs absorb oxygen normally and are histologically normal three months after operation, if the left pulmonary veins are divided and resutured as close as possible to the left atrium at the normal site of entry of these veins, provided a satisfactory atrial clamp and a careful evertting and locking suture technique are also used; (5) regarding the tolerance time of sheep lungs to anoxia, the left lung can regularly withstand two and a half hours of anoxia. There is no sharp end-point to tolerance of anoxia between three and four hours after circulatory arrest in the test lung. After four hours of anoxia with normothermia irreversible pulmonary oedema will regularly occur.

In lung transplantation in twin lambs, although the lungs functioned at first, the homograft reaction caused death from six to nine days after operation.

In lung transplantation in identical twin calves, the animals survived and had good initial post-operative air entry. Further, when on pasture they thrived. When, however, their test lungs were examined 55 and 48 days after operation, these were found to have been completely rejected by their host and to be encased in a fibrinous pleural cast.

From a biological viewpoint one should be able to transplant successfully identical twin calf lungs, but transplants of skin should also be done as well as relying on surface markings for establishing firm evidence of identical twinning. Immunosuppressants should improve these results.

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