Isolated Perfused Rabbit Lung: A Critical Appraisal

by Richard W. Niemeier*

The isolated perfused lung (IPL), when compared to available in vitro and in vivo pulmonary systems, is a preparation that fulfills a majority of the ideal criteria for studying metabolism, binding and/or physiological response to xenobiotics. The IPL is an exceptionally useful method when there is a need for concurrent administration of multiple agents in different physical forms. Various details such as physiological and biochemical parameters and the construction of a small animal tracheal valve system are discussed.

Our general interest in the lung originated from the fact that the respiratory tract is the main portal of entry and one of the first surfaces contacted by airborne contaminants. The main reason for our interest in pulmonary disposition of pollutants is the potential importance in the ultimate toxicity of some of these agents. Of interest, also, are the agents, drugs, or pollutants reaching the lung via the circulatory system. It has been well established that the lungs are capable of binding and/or metabolizing several such agents (1-12).

There is no way to study the pulmonary metabolic activity in vivo because of the influence of other organs. In vitro tissue preparations such as slices and homogenates compromise the integrity of an investigation (13), especially when considering concurrent administration of multiple agents in different physical forms or when determining distribution or binding of compounds throughout the pulmonary system. Therefore, the obvious choice in our opinion was the isolated perfused lung (IPL).

A number of criteria were chosen and considered mandatory in order to provide an isolated perfused lung preparation that was sufficiently stable to permit evaluation of metabolic activity, distribution, and uptake of compounds (14). In addition, we thought that monitoring of physiological and biological indices would better define the stability of the system. A summary of the major features of our isolated perfused rabbit lung preparation is presented in Table 1.

Table 1. Major features of isolated perfused lung preparation.

1. Perfusate: recirculation of undiluted, heparinized, autologous whole blood.
2. Constant blood pressure and blood flow
3. Chemically and biochemically inert
4. Normothermic conditions
5. Ventilation
   Subatmospheric pressure (-3 to -13 cm H2O)
   Net subatmospheric end respiratory pressure
   Sighing (-30 cm H2O)
   Fresh filtered gas, humidified and warmed
6. Monitoring of biochemical and physiological conditions
7. Controlled pH
8. Materials available for analyses
   Blood
   Lung washings
   Pulmonary tissues
   Ventilating gases

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Undiluted, heparinized, autologous whole blood was chosen as the perfusate for a number of reasons. The principal reason is that this perfusate is perhaps the best physiological and biochemical medium available, i.e., the essential cofactors, trace metals, and autologous proteins are present.

Investigations of benzo[a]pyrene metabolism have indicated that the metabolites are distributed differentially when comparing plasma to red cells (15). This observation reflects the importance of using whole blood when possible, since distribution, absorption, and excretion kinetics are important parameters in estimating total toxicity of a chemical. The possibility exists where significant factors may be overlooked when organs are perfused with artificial media. However, the design of the experiment may dictate the use of artificial media as for example in the study of lipid metabolism.

Figure 1 is a simplified schematic of our isolated perfused lung preparation (14) showing the lungs suspended in the artificial thorax. A peristaltic pump maintains the constant pressure of blood through an electronic feedback-level sensing device. Filtered air is warmed and humidified prior to passing the respiratory valve complex.

A summary of the biochemical changes found in the plasma of eight control isolated perfused rabbit lungs has been reported previously (14). One of the most notable changes is the glucose concentration changes, which are shown in Figure 2.

Table 2. Physiological values in the isolated perfused rabbit lung preparation.

| Parameter       | Value               |
|-----------------|---------------------|
| Hematocrit, %   | 35.0 ±5.0           |
| Mean change per hr | −1.6 ±0.3           |
| Weight gain, %/hr | 2.81 ±1.36          |
| Blood flow, ml/min | 180-240 (constant in each experiment) |
| $P_{O_2}$, mm Hg | 118 ±6, 121 ±10*    |
| $P_{CO_2}$, mm Hg | 39 ±4, 32 ±4*       |
| pH range        | 7.38-7.42           |
| Tidal volume, ml | 11.7 ±0.3, 11.0 ±0.4* |

* Typical values.
The average disappearance rate was approximately 35 mg-%/hr. Infusion of this amount resulted in no net change throughout the perfusion. Lactate dehydrogenase, glutamic oxalacetic transaminase (GOT), and lactic acid were found to increase quite substantially. The increases in LDH and GOT and the increases in plasma hemoglobin were attributed to hemolysis.

Typical physiological values obtained from control lungs are shown in Table 2. Historically, edema has been a problem in the IPL. We feel that we have minimized or at least delayed its appearance with this system, as is evident by the small increases in weight of the lungs measured before and after perfusion.

Cervical dislocation is used to kill the animal since anesthesia is undesirable for metabolic studies. The location of the strike must be precise, since brain trauma results in massive hemorrhage and edema in the lungs. We have also noted that in experiments in which the blood flow decreases with time in the IPL, edema and hemorrhage usually follows. Therefore, we sought to correct the blood flow problems that normally are found with the IPL hoping that these corrections might also influence edema formation.

Figure 2 shows typical blood flow values which we considered inadequate for metabolic studies before the corrective measures were made. It was evident that in 1-2 hr a consistent phenomenon was occurring. Areas of hemorrhage and low perfusion were very evident in these lungs. Administration of regular maintenance doses of heparin (200-500 IU) as well as epinephrine (40-100 μg) were found necessary to maintain constant blood flow in the isolated perfused rabbit lung (Fig. 3). Figure 4 illustrates the effects of...
blood flow of benzo[a]pyrene (BaP) in an ethanol saline (1:1, v/v) suspension administered intratracheally to the IPL. The initial decrease in flow is due to the ethanol administration. Histopathological examination of control lungs revealed no edema and excellent maintenance of pulmonary structures after 4 hr of perfusion.

We are primarily interested in the pulmonary metabolism of the ubiquitous carcinogen benzo[a]pyrene. One of our aims was to study this compound when coadministered intratracheally with ferric oxide or SO2. A development which arose from this need was the tracheal valve system which is shown in Figure 5. The valve is fabricated with Teflon and has an extratracheal dead air space of approximately 6 cm3. Silicone rubber stem valves permit unidirectional flow. The offset diagram in Figure 5 gives the dimensions of the valve extension mold which is also fabricated with Teflon. Intratracheal pressures can be measured and intratracheal instillations are made through a port at the top of the valve. Spirometric measurements are also possible, therefore, adding another dimension to metabolic and acute toxicity investigations.

Table 4. Pertubations with IPL.

| Perturbations prior to perfusion | Concurrent administration of multiple agents to IPL |
|---------------------------------|-----------------------------------------------|
| Enzyme-inducing agents (IP) Pb, 3-MC, B(a)P, PCBs | CAP + BaP ± SO2 |
| Inhalation exposure SO2, n-dodecane, coal dust, metals | Ferric oxide + BaP |
| Dietary manipulations | Ethanol + trichloroethylene |
| Intratracheal instillations | |
| BaP, crystalline quartz, papain, asbestos | |
| ferric oxide, with or without BaP | |
| crude air particulate (CAP), with or without BaP | |
| | |

The Rf values of the metabolites of benzo[a]pyrene found in isolated perfused rabbit lung preparation after intratracheal administration of 14C-BaP in ethanolic saline are listed in Table 3. The benene-reconstituted extracts were chromatographed on silica gel thin layer plates using benzene:ethanol (19:1). The Mylar-backed chromatograms were cut in 1-cm strips and placed in scintillation vials for subsequent counting.

Blood samples taken at various times during the perfusion were analyzed for their metabolite content. Typical values found in control lungs and those pretreated 24 hr prior to perfusion with 3-methylcholanthrene (3-MC) in corn oil (IP, 20 mg/kg) are shown in Figure 6. The rates of metabolic appearance are linear for 60 min or longer, and pretreatment with 3-MC was found to increase the total rate approximately sevenfold.

A summary of the pertubations that we have completed or are planning are presented in Table 4. This table reflects some of the potential uses of the IPL in characterizing the effects of many environmental contaminants on pulmonary metabolic activity. In addition, concurrent administration of multiple agents are made possible with this system for the purpose of investigating combined effects of agents in different physical forms.

### Table 3. Metabolites of BaP found in the IPL.

| Structure | Compound | Rf value |
|-----------|----------|----------|
| ![BaP](image) | BaP | 0.88 |
| ![BaP-3,6-dione](image) | BaP-3,6-dione | 0.78 |
| ![BaP-3-hydroxy](image) | BaP-3-hydroxy | 0.48 |
| ![BaP-4,5-dihydrodiol](image) | BaP-4,5-dihydrodiol (P1)* | 0.24 |
| ![BaP-7,8-dihydrodiol](image) | BaP-7,8-dihydrodiol (P1)* | 0.16 |
| ![BaP-9,10-dihydrodiol](image) | BaP-9,10-dihydrodiol (baseline)* | 0.03 |
| polar (conjugates)* | | |

* Tentative identification.
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