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Convective Heat Transfer within Mechanically-ventilated Building Spaces

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This thesis is dedicated

To the memory of my late father Dr. A. Alamdari

and

To my wife Fariba
A hierarchy of interacting and interdependent approaches have been developed for calculating internal surface convective heat transfer coefficients within mechanically-ventilated rooms. A 'high-level' computer code is developed for non-buoyant and buoyant flow based on the "elliptic" code of Pun and Spalding (1977), in which "upwind" finite-difference approximations to the governing partial-differential equations for continuity, momentum and thermal energy are formulated in terms of "primitive" pressure-velocity variables. Closure of these time-averaged, elliptic equations is obtained via transport equations for both the turbulence kinetic energy and its dissipation rate. The high-level code solves the difference equations for a predetermined size, staggered grid in an iterative "line-by-line" manner using a guess-and-correct procedure.

An "intermediate-level" computer code (the ROOM-CHT program) has also been developed for the above purpose, which employs "informed" estimates of the flow and thermal field based on the known mean flow properties of wall-jets. The corresponding heat transfer distribution across the room surface is calculated using wall-jet profile analysis or improved data correlations for buoyancy-driven convection as appropriate.

Computations are presented for a room into which air is injected through a low or high side wall register. The supply of air governed by both cyclic and modulating control was examined. The intermediate-level code is advocated as being the most appropriate for meeting the requirements of dynamic building thermal models. This code was verified by comparison with the high-level code and with experimental measurements. The computed heat transfer coefficients from the intermediate-level code were found to be in good agreement with that of the high-level code. Both indicate significantly higher values than those which would be obtained from established design guides. These high values suggest errors in building thermal models based on guide data, including substantial under-estimation of preheat times.
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NOTATION

a Coefficients in finite-difference equations, Eq.(3.17)

$a_1, a_2, a_3, a_4$ Coefficients in quatric wake function equation, Eq.(2.25)

$a_c, b_c$ Empirical coefficients in buoyancy-driven correlating equation, Eq.(4.37)

$a_i, b_i, c_i, d_i$ Coefficients in finite-difference (TDMA) for block-correction equation, Eq.(3.64)

A Area of the control volume face (m²)

Ag Grille area (m²)

$A_j, B_j, C_j$ Coefficients in finite-difference equations (TDMA), Eq.(3.58)

$A'_j, B'_j$ Transformed Coefficients (TDMA), Eq.(3.60)

$A_s$ Surface area (m²)

b The source term in finite-difference equation, Eq.(3.17)

$b_u, b_\theta$ 2D wall-jet velocity and temperature half width respectively (m), Eq.(4.1,4.8)

$b_z$ 3D wall-jet spanwise half width (m), Eq.(4.25)

$B, B_\theta, B', B'_\theta$ Log-law constants, Eq.(4.1,4.8,4.14)

$C_1, C_2, C_D, C_\mu, C'$ Coefficients in turbulence models, Eq.(2.17-2.21)

$C_f$ Skin friction coefficient, Eq.(2.24)

$C, C_\theta$ Velocity and temperature wake function coefficients, Eq.(4.1,4.8)

d Coefficient of pressure-difference term, Eq.(3.43)

D Diffusion terms, Eq.(3.19)

$D, D_\theta$ Wall-jet velocity and temperature width ratio, Eq.(4.3,4.10)
F Mass flow rate through a control volume (Kg/s), Eq.(3.14)

g Gravitational acceleration (m/s²), Eq.(2.41)

G_B Buoyancy production, Eq.(2.41)

G_k Generation rate of turbulence energy, Eq.(2.16)

h Instantaneous specific enthalpy (J/Kg),, Eq.(2.3)

h_c Convective heat transfer coefficient (W/ m²K), Eq.(4.35)

H Time-averaged enthalpy (J/Kg), Eq.(2.31), also room height (m)

H_g Grille height (m)

J Total (convection+diffusion) flux, Eq.(3.9), also diffusion flux Eq.(2.6)

K Turbulent kinetic energy, Eq.(2.15)

l_m Mixing length (m), Eq.(2.14)

l L Length scale, Eq.(2.17), also room length (m)

L_s Surface length (m), Eq.(4.38)

L_ε Dissipation length scale (m), Eq.(2.29)

N Number of air changed per hour (ACH), Eq.(5.2)

P Instantaneous static pressure (bar), Eq.(2.2)

P∗ Pressure-correction term (bar), Eq.(3.39)

P_ Time-averaged static pressure (bar), Eq.(3.39)

P^* Gussed pressure (bar), Eq.(3.39)

P_e Peclet number, Eq.(3.20)

P_s Surface perimeter (m)

q Heat flux (W/m²), Eq.(2.36)

Q Total heat load (W), Eq.(5.3)
R  Residual source, Eq. (3.69)
Re  Reynolds number at inlet
Rm  Maximum Reynolds number, Eq. (2.26)
Sb  Buoyancy source term, Eq. (2.10)
Si  Slot inlet height (m), Eq. (2.31)
Sp  Constant part of linearized source term, Eq. (3.11)
Sp  Coefficient of $\phi_p$ in linearized source expression, Eq. (3.11)
Sf  General source term, Eq. (3.11)
St  Stanton number, Eq. (4.12)
t  Time (s), Eq. (2.1)
T  Temperature (K), Eq. (4.8)
Tf  "Friction temperature" (K), Eq. (4.8)
u, u1  x-direction velocity component (m/s), Eq. (2.1)
$u^*$, $u_1^*$  Fluctuating component of x-direction velocity (m/s), Eq. (2.10)
U, U1  Time-averaged x-direction velocity (m/s), Eq. (2.9)
U*  x-direction velocity based on the gussed pressure $P^*$ (m/s), Eq. (3.38)
U  x-direction pseudo-velocity (m/s), Chapter 6
Um, Umo  Maximum and centreline maximum velocity respectively (m/s), Eq. (2.24, 4.24)
Ut  "Friction velocity" (m/s), Eq. (4.2)
v, u2  y-direction velocity component (m/s), Eq. (2.1)
v*, u_2*  Fluctuating component of y-direction velocity (m/s), Eq. (2.10)
V, U2  Time-averaged y-direction velocity component (m/s), Eq. (2.9)
V*  y-direction velocity based on the gussed pressure $P^*$ (m/s), Eq. (3.38)
\( \hat{V} \) \hspace{1cm} y-direction pseudo-velocity (m/s), Chapter 6

\( \dot{V} \) \hspace{1cm} Volume flow rate (m\(^3\)/s), Eq. (5.3)

\( W \) \hspace{1cm} Weighting function, Eq. (3.4), also room width (m)

\( W_{gr} \) \hspace{1cm} Grille width (m)

\( x, y, z \) \hspace{1cm} Cartesian coordinates

\( x_{c} \) \hspace{1cm} Potential core length (m), Eq. (4.31)

\( X_{sp}, Y_{sp} \) \hspace{1cm} Location and width of the starting profiles (m), Eq. (2.23)

\( y^{+} \) \hspace{1cm} Dimensionless distance from the wall, Eq. (4.1)

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**Greek Symbols**

\( \alpha \) \hspace{1cm} A large number (for example 10\(^{30}\)), Eq. (3.57)

\( \beta \) \hspace{1cm} Coefficient of thermal expansion, Eq. (2.41)

\( \Gamma \) \hspace{1cm} Diffusion coefficient, Eq. (2.40)

\( \delta \) \hspace{1cm} Wall-jet velocity or temperature width, Eq. (2.28), and also the distance between near boundary and boundary node (m), Eq. (3.49)

\( \delta x, \delta y \) \hspace{1cm} Inter-nodal distances, Eq. (3.19)

\( \Delta x, \Delta y \) \hspace{1cm} Dimensions of the computational cell, Eq. (3.10)

\( \varepsilon \) \hspace{1cm} Rate of dissipation of K, Eq. (2.17)

\( \kappa - \varepsilon \) \hspace{1cm} Log-law constants, Eq. (2.34)

\( \theta \) \hspace{1cm} Excess temperature (K), Eq. (2.41)

\( \lambda \) \hspace{1cm} Convergence criteria, Eq. (3.70)

\( \mu \) \hspace{1cm} Fluid dynamic viscosity (Kg/ms), Eq. (2.5)

\( \nu \) \hspace{1cm} Fluid kinematic viscosity (m\(^2\)/s), Eq. (4.2)

\( \xi \) \hspace{1cm} Relaxation factor, Eq. (3.73)
ρ Fluid density (Kg/m^3), Eq. (2.1)
σ Prandtl/Schmidt number, eq. (2.6)
τ Stress-tensor, Eq. (2.5)
τ_w Wall shear-stress, Eq. (2.34)
φ Instantaneous general scalar property, Eq. (2.4)
φ' Fluctuating value of φ, Eq. (2.8)
φ̅ Time-averaged value of φ, Eq. (2.7)

Subscripts

a Ambient condition
e Inlet condition
eff Effective value
e, w, n, s Control volume faces
P, E, W, N, S Central node and its neighbours
ij Components in cartesian tensor notation
i, j Location of point in cartesian grid
l Laminar flow condition
m Maximum value
o Centreline and also reference value
r Room condition
ref Reference value
t Turbulent flow condition
w Condition at wall
1,2 Cartesian directions
Superscripts

+  Dimensionless value
-  Average value
~  Approximate solution
^  Pseudo-velocities
*  Previous iteration value, also velocities based on guessed pressure
  Correction to starred values, also fluctuations relative to the time-averaged values

Special Symbol

Max (A, B)  The bigger of A and B
PART I

INTRODUCTION
CHAPTER 1
ENERGY CONSERVATION AND THE BUILT ENVIRONMENT:

1.1 DYNAMIC THERMAL PERFORMANCE OF BUILDING:

It has become apparent, following the "energy crisis" of the mid-1970's, that the world's reserves of fossil fuel are exhaustible. Consequently, it is necessary to make the most effective use of energy, thereby extending the available time for developing alternative energy supplies. The world consumption is accelerating towards the energy "bankruptcy" (see Fig 1.1), instead of reducing or at least maintaining steady-state. In many countries the energy used in domestic and commercial buildings is a large proportion of total energy consumption. Most energy consumed in buildings is used for space heating, cooling and ventilation, so that saving in these areas are of great importance.

In the last decade it has been become increasingly recognised that in order to develop realistic methods for the energy-conscious design of buildings it is necessary to model the dynamic thermal response of the system. These dynamic models normally require computational solution (as noted in the reviews by Clarke (ref 1) and Day (ref 2); the latter summarising the efforts of the U.K Science and Engineering Research Council to stimulate research in this area), in contrast to the manual calculation methods used with steady-state procedures. However, progress in this direction has been hindered by uncertainties in some of the input data, particularly ventilation rates and surface heat transfer coefficients.

The convective heat transfer data currently available to the designer is certainly inadequate for modern requirements. ASHRAE (ref 3) seem to ignore the possibility of forced convective heating, and employ only buoyancy-driven convective internal surface coefficients to obtain fabric "U-values". The recent edition of the CIBS guide (ref 4) provides an approximate correction factor to buoyancy-driven data when air velocity over individual surfaces has non-zero values. Unfortunately this practice is not very helpful as the designer generally has no means of determining this velocity a priori. In any case, such a procedure neglects variations in surface coefficients due to different heating system/room configurations. In a field study of over thirty rooms of various shapes and sizes heated by a fan "convector", Yaneske and Forrest (ref 5) found the room-averaged convection coefficient to be 6.31 W/m²K, but with a wide scatter: the standard deviation of their data was 4.77 W/m²K. They also showed that, when selecting heater capacity, the use of a surface coefficient appropriate to buoyancy-driven convection (3.00 W/m²K), instead of a forced value, would result in substantial increase in preheat times. More recently, Waters (ref 6) has similarly shown that the accuracy of building thermal models for
mechanically-ventilated rooms are strongly dependent on the choice of convection coefficient. He reached this conclusion using an implicit finite-difference model having a relatively fine time-space grid. The time step adopted was only about 5 minutes, in contrast to the 60 minute steps typically employed in response factor programs. Irving (ref 7) has noted, when reviewing the achievements of the International Energy Agency (IEA) Annex 1 validation studies, that uncertainties in input data currently limit the level of sophistication to which it is worthwhile developing the overall building thermal model.

Warm air heating systems, probably incorporating a ventilation capability, have many potential advantages for low energy housing of the future. These benefits include a good energy efficiency (70-75% over heating season), and ease of control with rapid response to load changes. In practice, the operation of the circulating fan in conventional warm air heating systems is controlled by an on-off thermostat, and therefore the warm-air is injected into room intermittently (ref 8-9). The rate of heat transfer at room internal surfaces will alternate between those corresponding to forced and buoyancy-driven convection. An alternative modulating control system may be utilised, in which a variable-speed fan is operated continuously, except at low load when cyclic operation again occurs. Pimbert (ref 8) has shown, on the basis of laboratory tests using a "Comfy-test" meter, that modulating control results in an improved thermal environment at the most frequently experienced loads. The room heat transfer coefficient under this control system correspond to forced convection, although their magnitude will vary with fan speed and supply air temperature as the heating load changes.

1.2 ROOM AIR MOVEMENT AND THERMAL COMFORT:

The air-movement within mechanically ventilated enclosures is a complex, jet-induced turbulent flow which gives rise to a recirculation pattern and therefore uncertainty in thermal comfort variables such as velocity, pressure and temperature distribution. Human discomfort could arise due to draught sensation if the air velocity exceed about 0.15 to 0.24 m/s depends on the prevailing temperature (ref 10-17). Conversely a feeling of stuffiness may be caused if the air velocity falls below about 0.05 m/s (ref 8-9).

In warm air systems, discomfort may arise due to the re-establishment of infiltration draughts during the "off" period of the control cycle (ref 18). Indeed this discomfort has inhibited the introduction of warm-air systems despite their potential for "low energy housing". The influence of this intermittent operation on the character of the injected jet and subsequent room air-movement has received little attention in the published literature.
1.3 THE AIM OF THE PRESENT WORK:

The present work was originally aimed at developing a computer program which would be capable of predicting the convective heat transfer coefficient over the internal surface of two-dimensional, mechanically-ventilated enclosures. However, an "intermediate-level" calculation procedure for three-dimensional enclosures was subsequently developed. It was also hoped that such a program would facilitate a better understanding of physics of the flow generated by domestic warm-air heating systems. In this way the requirements of building thermal modellers for input heat transfer coefficients of known uncertainty would be met.

1.4 APPROPRIATE CALCULATION METHODS FOR THERMO-FLUID PROBLEMS RELATING TO BUILDINGS:

Thermo-fluid calculation methods can, according to Hammond (ref 19), be classified in one of the following categories:

a) "Lower-level" Methods

This category uses analytical solutions and empirical data for simple shear flows, and thus apply to a very narrow class of flows or range of calculations. An examples of this category are the profile analysis of wall-jets and improved data correlation for buoyancy-driven convection in rooms which are outlined in part III and appendix A of the present work.

b) "Higher-level" Methods

The arguably most sophisticated approach is the development of "higher-level" computer-based flow models (one is described in detail in part II), which involve the solution of the governing equations of the flow. The model used for the present study used the "elliptic" finite-difference computer code of Professor Spalding and his co-workers, together with a two-equation turbulence closure. Unfortunately, such high-level flow models are unsuitable for providing direct input data into the current generation of building thermal models. This is because the models are very costly in terms of computer time and storage, and are rather user "unfriendly", due to their complexity.

c) "Intermediate-level" Method

In order to overcome the practical limitation of "higher-level" methods, an "intermediate-level" calculation procedure has been developed by Alamdari and Hammond (ref 20) and incorporated into a computer code called the ROOM-CHT (Room Convective Heat Transfer) program. This program was envisaged as a "bolt-on" package to building thermal models, and is outlined in part III of the present work.
1.5 LAYOUT OF THE THESIS;

The present work concentrates mainly on the development of "high-level" and "intermediate-level" computational methods for convection within mechanically-ventilated enclosures, and these are described in part II (Chapters 2 & 3) and part III (Chapter 4) of this thesis respectively.

Chapter 2 lays the mathematical foundation for the high-level approach, while Chapter 3 deals with the numerical solution procedure for the governing flow equations.

Chapter 4 explains the physical basis of the intermediate-level calculation method.

The computed heat transfer coefficients and the experimental validation of the predictions for both the high-level and intermediate-level methods are discussed in Chapter 5 (part IV). The results of both the two and three-dimensional versions of the intermediate-level method are compared. The results of a parametric studies using the two-dimensional version of the intermediate-level program for the case of commercial office air-conditioning via linear or slot "diffusers" are then presented. In the final section of Chapter 5 the sensitivity of building models to convection data is assessed. In the final chapter of this thesis (Chapter 6) general conclusions are drawn concerning the validity of the various calculation methods "levels", and recommendations are made for further research.
PART II

A HIGHER-LEVEL COMPUTER BASED MATHEMATICAL MODEL
CHAPTER 2
MATHEMATICAL FRAMEWORK FOR HIGHER-LEVEL MODEL

2.1 INTRODUCTION:

"Higher-level" mathematical models of the type employed here were developed in the early 1970's for predicting complex elliptic flows in practical configurations. These methods solve numerically the differential equations for continuity, momentum and thermal energy in order to generate field values for the velocity components, the static pressure and the enthalpy or temperature. The mathematical statement of the physical laws which govern the flows, and the way that the problem is formulated in terms of physical differential equations for laminar and turbulent flow regimes are briefly described in this Chapter.

2.2 OUTLINE OF THE CALCULATION PROCEDURE:

2.2.1 Previous Work:

In recent years there has been considerable progress in the application of numerical computation procedures for fluid flow and heat transfer problems, although relatively few studies have been related to ventilation and heat transfer problems in building spaces. However, in the mid-1970's Neilsen (ref 21) developed a model for the prediction of two-dimensional, isothermal flow in rooms, by using an early stream-function and vorticity procedure (ref 22), and employing the two-equation, K-ε model of turbulence (ref 23-24) in order to close the governing time-averaged equations. In this method the stream-function and vorticity act as a main dependent variables instead of the velocity components, thereby eliminating the pressure gradient terms which appear in the momentum equations. This method cannot be extended to three-dimensional flows, and the scheme has a poor convergence performance.

Neilsen et al (ref 25-27) have more recently solved the same problem as above but using the velocity components as dependent variables, thereby enabling extension to three-dimensional flows. Neilsen et al (ref 26) also attempted to model the effect of buoyancy, by introducing the buoyancy terms into the y-direction momentum, and turbulence energy and dissipation rate equations, through their source terms. They examined the case of cool-air injection with a predominant "through-flow", and relatively small buoyancy effects. The through-flow geometry is an idealisation which greatly assists convergence, but is rarely found in practice.
Hjertager and Magnussen (ref 28) have used the Patankar and Spalding (ref 29) "SIMPLE" algorithm in order to predict three-dimensional turbulent flow in a ventilated room for both isothermal and buoyant flows, again with the two-equation, K-ε model of turbulence. The influence of buoyancy was introduced into the problem only through the y-direction momentum source terms. A cooled jet (11 K lower than the averaged room temperature, Tr=293K) was emitted horizontally from a rectangular nozzle placed just under the ceiling with nearby outlets extracting the flow vertically to the ceiling cavity. Part of the floor and far-wall (the wall opposite to the jet) were heated in the case of buoyant flow, which is again rare in domestic and commercial office spaces in the UK.

Sakamoto and Matsuo (ref 30) also examined a three-dimensional turbulent flow in a ventilated room, but for isothermal flows only. It seems their aim was mainly to evaluate the use of different turbulence models. The geometry which they considered was a cubic room with a square nozzle inlet at the centre of the ceiling and outlet at the one of the walls closed to the floor.

2.2.2 The Present Work:

In the present work a modified version of the "elliptic" computer code of Pun and Spalding (ref 31) has been used as the basis for a "higher-level" flow model for mechanically-ventilated, "two-dimensional" rooms. In this code, "upwind" finite-difference approximations to the governing elliptic equations are formulated in terms of primitive pressure-velocity variables for a predetermined size, staggered grid. The SIMPLE (Semi-Implicit Method for Pressure-Linked Equations) algorithm (ref 29,32) was used in the code to solve the finite-difference equations in a "line-by-line" manner with nearly-exact adjustment of terms. The original continuity (mass conservation) adjustment technique used by Pun and Spalding (ref 31) was found to be unsuitable for the present case of severe flow recirculation together with prescribed jet inlet conditions. It was therefore replaced by a "block-correction" procedure, which also applied in a line-by-line manner, after each iteration. The jet inlet conditions were specified in a similar manner to that recommended by Nielsen et al (ref 25-26), but using wall-jet empirical data and results of profile analysis of Hammond (ref 33-34). The present work has concentrated on the problem of handling flows involving complete recirculation of the room air-flow which is a much more severe test of the model than the geometry studied by Neilson et al (ref 25-26). The present study also examined flows with significant buoyancy, which gives rise to additional problems of ensuring numerical stability and in turbulent modelling.

The treatment of turbulence is made by way of mathematical model (ref 23), in which the transport of momentum by the
turbulent motions is related to a turbulent viscosity, and transport of scalar quantities. The Launder & Spalding (ref 24) version of "K-ε" model has been used, in which the turbulent viscosity and diffusivity are evaluated from two local properties of turbulence, namely time-averaged kinetic energy "K" and dissipation rate "ε".

The buoyancy-influenced terms have been introduced into the y-direction momentum, and turbulence energy and dissipation rate equations in the similar to those adopted by Nielsen (ref 26), Markatos and Cox (ref 35) and Kumar and Cox (ref 36). The latter two studies concerned fire spread within an enclosure, which is driven by large buoyancy forces.

2.3 GOVERNING EQUATIONS AND THEIR FINITE-DIFFERENCE ANALOGUE:

The higher-level numerical solution procedure is based on the laws which governing the physics of the flow field in terms of partial differential equations (pdes), which govern the conservation of mass, momentum and thermal energy for any laminar and turbulent regime. The time-averaged pdes for a turbulent flow are the same as the laminar ones, except that effective transport properties are employed. The dependent variable of these equations are usually specific properties, and are based on a unit-volume.

2.3.1 The Continuity, Momentum and Energy Conservation Equations:

The conservation equations which govern the buoyancy-driven and forced convection of mass, momentum and thermal energy may be expressed in cartesian tensor notation as follows (ref 37-39):

a) Mass conservation or continuity equation:

$$\frac{\partial \rho}{\partial t} + \frac{\partial}{\partial x_j} (\rho u_j) = 0 \quad (2.1)$$
b) Momentum or Navier-Stokes equations for direction i:

\[
\frac{\partial}{\partial t} (\rho u_i) + \frac{\partial}{\partial x_j} (\rho u_i u_j) = \frac{\partial}{\partial x_j} (\tau_{ij}) - \frac{\partial p}{\partial x_i} + S_{b,i}
\]  

(2.2)

c) Thermal energy equation:

\[
\frac{\partial}{\partial t} (\rho h) + \frac{\partial}{\partial x_j} (\rho h u_j) = \frac{\partial}{\partial x_j} (J_{h,j}) + S_h
\]  

(2.3)

where

- \( i \) & \( j \) are components in cartesian tensor notation, both can take the values of 1 and 2 denoting the directions \( x_1 \) and \( x_2 \) in two space coordinates. When a subscript is repeated in a term, a summation of two terms is implied,

- \( x_1 \) & \( x_2 \) are \( x \) and \( y \) components in cartesian coordinate,

- \( \rho \) is the fluid density,

- \( t \) is the time,

- \( u \) is the instantaneous velocity,

- \( p \) is the instantaneous static pressure,

- \( S_b \) is buoyancy force,

- \( \tau \) is the stress-tensor (see Section 2.3.3),

- \( h \) is the instantaneous specific enthalpy,

- \( J_{h} \) is the heat-flux tensor and

- \( S_h \) is the source/sink of \( h \) (negligible unless chemical reaction is present).

Equations (2.1), (2.2) and (2.3) are in fact a closed set of equations which must be solved simultaneously, by a finite-difference method in the present study, in order to obtain the velocity, pressure and enthalpy in the field.
2.3.2 The Conservation Equation For A General Scalar Property:

It is convenient to write a conservation or transport equation for a general scalar property of the flow such as enthalpy/temperature. It will be shown later that the turbulence kinetic energy $K'$ and its dissipation rate $\epsilon'$, have similar transport equations, since they obey the generalized conservation principle. If the dependent variable is denoted by $\phi$, the general differential equation is:

$$\frac{\partial}{\partial t} (\rho \phi) + \frac{\partial}{\partial x_j} (\rho \phi u_j) = \frac{\partial}{\partial x_j} (J_{\phi,j}) + S_{\phi}$$

(2.4)

where $S_{\phi}$ is the source/sink of $\phi$ and $J_{\phi,j}$ is the scalar property flux.

2.3.3 The Auxiliary Relations:

The molecular diffusion of momentum, $\tau_{ij}$, and the scalar property flux, $J_{\phi,j}$, in equations (2.2) and (2.4) have not yet been defined. They can be related to gradient of velocity components and scalar properties respectively. For uniform density flows these relations may be written as follows (ref 22).

$$\tau_{ij} = \mu \left( \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} \right)$$

(2.5)

$$J_{\phi,j} = \frac{\mu}{\sigma_{\phi}} \left( \frac{\partial \phi}{\partial x_j} \right)$$

(2.6)

where $\mu$ and $\sigma_{\phi}$ are the dynamic viscosity and molecular Prandtl/Schmidt number of the fluid respectively.

2.4 Time-Averaged Equation for Turbulent Flow:

The calculation of turbulent flows via the exact time-dependent forms of the governing equations is possible, although it is limited to very low Reynolds numbers and requires an excessively fine grid, which renders this approach uneconomic (ref 40). It is therefore necessary to utilise the time-averaged version of these equations, in which the instantaneous properties of the flow are replaced by mean values with fluctuating values.
superimposed upon them (ref 40, 41). For a general property or variable, \( \phi \) (ie. \( u, p, h, \) etc. . . .):

\[
\phi = \frac{1}{t} \int_{0}^{t} \phi \, dt \quad \text{as} \quad t \to \infty
\]

(2.7)

\[
\phi = \phi + \phi'
\]

(2.8)

where \( \phi \) and \( \phi' \) are the time-averaged component and fluctuating component of the variable respectively.

By substituting equation (2.7) into equations (2.1), (2.2) and (2.4), and thereafter time-averaging (ref 42), the following differential equations will result which govern the mean motion of "steady" (strictly statistically stationary) turbulent flow:

a) **Continuity:**

\[
\frac{\partial}{\partial x_j} (\rho \, U_i) = 0
\]

(2.9)

b) **Momentum:**

\[
\frac{\partial}{\partial x_j} (\rho U_i \, U_j) = \frac{\partial}{\partial x_j} \left\{ \mu \left( \frac{\partial U_i}{\partial x_j} + \frac{\partial U_j}{\partial x_i} \right) - \rho U_i' U_j' \right\}
- \frac{\partial \rho}{\partial x_i} + S_{b,i}
\]

(2.10)

c) **General Scalar Property:**

\[
\frac{\partial}{\partial x_j} (\rho \phi \, U_j) = \frac{\partial}{\partial x_j} \left\{ \frac{\mu}{\phi} \left( \frac{\partial \phi}{\partial x_j} \right) - \rho \phi' U_j' \right\} + S_{\phi}
\]

(2.11)

where \( U \) is the time-averaged velocity,
is the time-averaged static pressure,
\( \phi \) is the time-averaged general scalar property,
\( S_b \) is the time-averaged buoyancy force,
\( S_\phi \) is the time-averaged source/sink of \( \phi \),
\( u' \) is the fluctuating value of \( u \),
\( p' \) is the fluctuating value of \( p \),
\( \phi' \) is the fluctuating value of \( \phi \),

and \( \tau \) & \( J_\phi \) have been replaced with relations (2.5) and (2.6) respectively.

The "Reynolds" stress-tensor (-\( \rho u' u'^j \)) and heat-flux tensor (-\( \rho \phi' u'^j \)) are unknowns and therefore before any solution can be obtained they must be expressed in terms of known or calculable quantities.

2.5 TURBULENCE MODELS:

The task of expressing the unknown fluxes (-\( \rho u' u'^j \), -\( \rho \phi' u'^j \)) in equations (2.10) and (2.11) in terms of the mean properties of the flow is known as the turbulence closure problem. Mathematical expressions which perform this function are called "turbulence models".

The applicability, economy and accuracy of a turbulence model are the main factors which determine its choice. Many turbulence models are available, and some of the more popular ones are described briefly in this Section. However, a more detailed description of them can be found in (ref 23), (ref 41-45) or elsewhere. There is no ideal classification scheme for turbulence models, but the one suggested by Reynolds (ref 41), and used below, provides a systematic basis for describing the various models.

2.5.1 The Classification of Turbulence Models:

a) Zero-equation models:

The zero-equation models are those in which no partial differential, or transport, equations for turbulence quantities are used. An example of such models is the classical Prandtl mixing length hypothesis, which employs the Boussinesq eddy viscosity formulation (ref 45). In this approach, the turbulent transport is related to the gradients of mean flow quantities, for example in thin shear layers:
\[- \rho \ u_1 \ u_2 = \nu_t \left( \frac{\partial U}{\partial x_2} \right) \]  

(2.12)

\[- \rho \ u_2 \ \phi = \frac{\nu_t}{\sigma_t} \left( \frac{\partial \phi}{\partial x_2} \right) \]  

(2.13)

where $\nu_t$ and $\sigma_t$ are the turbulent viscosity and Prandtl/ Schmidt number respectively.

The Prandtl mixing length hypothesis calculate the distribution of the turbulent viscosity, $\nu_t$, by relating it to the local mean velocity gradient:

$$\nu_t = l_m^2 \left| \frac{\partial U}{\partial x_2} \right|$$  

(2.14)

where $l_m$ is the mixing length, and must be described over the flow field empirically.

b) **One-equation model**

These models use one partial differential equation (pde) relating to some turbulence velocity scale. Usually the scale is the square root of the kinetic energy of turbulence, $\sqrt{K}$. The modelling proposal of Launder and Spalding (ref 23) and others lead to the following equation for $K$, valid for high Reynolds number flows (i.e. when the effect of molecular viscosity can be neglected).

$$\frac{3}{\partial x_j} \left( \rho K \ u_j \right) = \frac{3}{\partial x_j} \left( \frac{\nu_t}{\sigma_k} \frac{\partial K}{\partial x_j} \right) + G_k - \rho \varepsilon + S_{b,k}$$  

(2.15)

and

$$G_k = \nu_t \left( \frac{\partial U_i}{\partial x_j} \left( \frac{\partial U_i}{\partial x_j} + \frac{\partial U_j}{\partial x_i} \right) \right)$$  

(2.16)

where

* $\varepsilon$ is the dissipation rate,
* $\sigma_k$ is Prandtl/Schmidt number for $K$ and
* $G_k$ is the generation term.
In these models, the dissipation rate is usually determined from:

\[ \varepsilon = C' \frac{K^{3/2}}{L} \]  

(2.17)

where

- \( L \) is length scale and must be prescribed via empirical information similar to that for \( l_m \), and
- \( C' \) is a constant value.

The eddy viscosity concept can be used here to relate the turbulence stresses to the turbulence kinetic energy determined from equation 2.14. This approach yields the so-called Kolmogorov-Prandtl expression (ref 45):

\[ \mu_t = C_\mu \rho K^{3/4} L \]  

(2.18)

where

- \( C_\mu \) is a constant

Bradshaw et al (ref 46) developed a popular one-equation model without using the eddy-viscosity concept. They have converted the turbulence kinetic energy equation into a transport equation for the shear stress \((\rho u_i u'_j)\) by assuming a direct link between \( K \) and \( \rho u_i u'_j \). They also assumed the diffusion flux of \( K \) to be proportional to a bulk velocity.

c) Two-equation models

Two-equation models employed pde's for both the turbulence velocity and length scales, this overcomes the need to specify the length scale empirically, which is difficult in complex, recirculating flows. However, the length scale itself is not normally used as the dependent variable in a pde. One successful model used the turbulence energy dissipation rate in place of the length scale equation, and is thus known as \( K-\varepsilon \) model of turbulence. In this model the Reynolds stress \((-\rho u'_i u'_j)\) is again related to mean velocity via an eddy viscosity which is calculated from the kinetic energy, \( K \), and its dissipation rate, \( \varepsilon \), via the Kolmogorov-Prandtl expression.
\begin{equation}
- \rho u_i u_j = \mu_t \left( \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} \right)
\end{equation}

\begin{equation}
\mu_t = C_D \rho^2 K^2 / \varepsilon
\end{equation}

where \( C_D \) is an empirical constant.

The transport equation for the dissipation rate, \( \varepsilon \), for high Reynolds numbers is as following (ref 23),

\begin{equation}
\frac{\partial}{\partial x_j} (\rho \varepsilon U_j) = \frac{\partial}{\partial x_j} \left( \frac{\mu_t}{\sigma_\varepsilon} \frac{\partial \varepsilon}{\partial x_j} \right) + C_1 G_k \frac{\varepsilon}{K} \\
- C_2 \rho \frac{\varepsilon^2}{K} + S_{b,\varepsilon}
\end{equation}

where \( C_1 \) and \( C_2 \) are empirical constants, and \( \sigma_\varepsilon \) is an effective Prandtl / Schmidt number.

d) Stress-equation models

Models which are based on eddy-viscosity/diffusivity concept such as the \( K-\varepsilon \) model, are valid only under certain conditions. In these models the eddy viscosity is assumed to be the same for all turbulent stresses, although in complex flows or flows with turbulence influenced by strong body forces acting in preferred directions, such as buoyancy forces, it may be anisotropic. The local state of turbulence is also characterized in these models by one velocity scale, \( \sqrt{K} \), and individual stresses are related to this scale. However, in reality, each stress component may develop quite differently in the flow.

In order to account for the different development of individual stress components pde's for all components of turbulence stresses are necessary. These models are often called second-order closure schemes, and a variety of such models have been proposed by different investigators (ref 44,47-49). The pde's for these models are complex and lengthy, and have the general form as follows:
D(\text{du}_i\text{du}_j)/Dt = \text{Production} + \text{Dissipation} + \text{Redistribution} \quad (2.22)

e) Large-eddy simulations models:

In 1973 the group at Stanford University under Professor W. C. Reynolds with close cooperation from the NASA-Ames laboratory began to develop this approach, and considerable success has been achieved since that time. However, the method is not able to be used for complex problems, but it might after considerable development, eventually be useful to handle problems as such.

The basic idea of this approach is that a numerical computation of the time-dependent Navier-Stokes equations is performed for the large-eddy structure while a sub-grid effective viscosity is used for the small scales turbulence. A more detailed description of the progress in large-eddy simulation is given by Reynolds (ref 41).

2.5.2 The Turbulence Model Adopted For Present Study:

Zero-equation models have been used widely for simple shear layer problems, where the upstream "history" of turbulent transport is not important and a mixing length distribution can be prescribe realistically. In one-equation models a pde for turbulent kinetic energy is solved, and thus these transport effects can be taken into account. However, the problem of specifying the length scale distribution in complex flows remains. Therefore, zero-equation and one-equation models are not suitable for the present work, where the flow patterns are complex and description of mixing length or length scale is difficult.

The stress-transport models appear to be potentially more accurate and "universal", but they require much more computer time than two-equation models, since there are more pdes to be solved. The validation studies that have compared the two-equation models with stress-transport models, such as that by Pope and Whitelaw (ref 50), have shown that the predictions of the latter models display little improvement. The two-equation model was therefore adopted for the present study simply because it is capable of handling complex recirculating flows without the need to prescribe any length scale, and it uses much less computer time than the stress-transport models.

The present work has concentrated on "K-\varepsilon", two-equation model of turbulence since it has been tested widely in many different flows and found to have an accuracy sufficient for practical purposes. The values of the turbulence model constants (i.e. $C_1$, $C_2$, $C_D$, $\alpha_k$, $\alpha_c$) employed here are those adopted in the elliptic code of Pun and Spalding (ref 31), and are given in Table 2.1. The basis for the determination of these constants is
explained by Launder and Spalding (ref 23).

<table>
<thead>
<tr>
<th>Table 2.1; Constants in turbulence model</th>
</tr>
</thead>
<tbody>
<tr>
<td>C₁</td>
</tr>
<tr>
<td>1.43</td>
</tr>
</tbody>
</table>

2.6 BOUNDARY CONDITIONS:

A two-dimensional room (see Fig 2.1) is typically modelled in the present work with a low side "linear" slot wall register. Thus the solution domain is bounded by walls, the floor, the ceiling, and slot inlet and outlet planes.

The condition at any point in the flow can be influenced by other points, because of "elliptic" nature of the flow. Therefore it is necessary to supply boundary conditions for variables at all the boundaries of the flow domain, these boundaries could be specified value of the variable or value of the associated flux, or a relation between the variable and the flux. Since the velocities and pressures are inter-dependent, where velocities are prescribed pressure need not. Therefore the variables for which boundary conditions are supplied are U, V, H, K and ɛ.

The boundary conditions specified for the present study are as follows:

2.6.1 At Jet Inlet and Outlet:

Particular difficulties arises with the use of numerical finite-difference methods in a large room with a very small slot inlet, due to the large number of grid nodes that would be required in the slot. In the present study inlet conditions were specified in a similar manner to those recommended by Nielsen et al (ref 25), but using wall-jet empirical data and the results of profile analysis of Hammond (ref 33,34). In this way an arbitrary inlet and outlet have been prescribed in the fully developed region of the wall-jet (as shown in Fig 2.1). The location of the starting profile, Xsp, is unimportant, provided that is kept small compared to the length of the room. In the present calculation the distance Xsp is located at the nearest grid line to the 1/4 of the room length. The width of the starting zone, Ysp, can be then estimated from the growth of the jet (Hammond private communication, 1981):
where \( S_i \) is the height of the slot inlet. The actual value of the width \( Y_{sp} \) corresponds to distance to the next farthest grid line away from floor beyond that given by equation 2.23.

### 2.6.1.1 Starting profiles:

a) Velocity profile in \( x \)-direction

Using complete velocity profile formula and associated data correlations given by Hammond (ref 33,34), (see also part III) gives:

\[
U = U_m \left( \frac{C_f}{2} \right)^{\frac{1}{2}} \left\{ \frac{1}{k} \ln R_m \left( \frac{C_f}{2} \right)^{\frac{1}{2}} \left( \frac{b_u}{y_m} \right) \left( \frac{y}{b_u} \right) \right\} + B + C \omega \left( \frac{y}{b_u} \right)
\]

(2.24)

A quartic wake function was adopted:

\[
\omega \left( \frac{y}{b_u} \right) = \sum_{n=4}^{n_{max}} a_n \left( \frac{y}{b_u} \right)^n
\]

(2.25)

where

- \( U \) is the velocity component in \( x \)-direction,
- \( U_m \) is the maximum velocity,
- \( C_f \) is the local skin friction coefficient,
- \( y \) is the normal distance from the wall,
- \( b_u \) & \( y_m \) are defined in fig 2.1,
- \( B \) is the log-law constant (=5.2, see part III),
- \( C \) is the wake function coefficient (see part III)
Hammond (ref 34) presented approximate values for the coefficients, $a_n$, as power-law functions of the local, wall-jet Reynolds number ($R_m = \rho U_m y_m / \mu$):

\[
\begin{align*}
    a_1 &= 0.841 R_m^{-0.1040} \\
    a_2 &= 0.223 R_m^{0.0917} \\
    a_3 &= -0.18 R_m^{0.0627} \\
    a_4 &= 0.030 R_m^{0.0555}
\end{align*}
\]

(2.26)

These approximate relations are used to specify the $U$-component starting profile.

b) Velocity Profile in $y$-direction

It is assumed that the flow is fully developed and the velocity component in direction $y$ is zero:

\[ V=0 \]

(2.27)

c) Turbulence kinetic energy profile

Hammond (private communication, 1981) has derived an empirical expression for the turbulence kinetic energy profile across a wall-jet, which may be written in the form:

\[
K = \left\{ U k_1 \left[ k_2 \left( \frac{y}{\delta} \right)^2 + 1 \right] \right\}^2
\]

(2.28)

where $\delta$ is the velocity width and the values of $k_1$ and $k_2$ are 0.175 and 7.10 respectively.

d) Energy dissipation profile

Energy dissipation profile for wall-jet may be obtained via the mixing length hypothesis. Hammond (private communication, 1981) has derived a dissipation length scale expression for plane wall-jets in the form:
where \( L_{e} \) is the dissipation length scale. The dissipation rate may then be obtained from the definition:

\[
\varepsilon = \frac{k^{3/2}}{L_{e}}
\]  

(2.30)

e) Enthalpy/ Temperature profile

The rather limited data for heated wall-jets on an adiabatic surfaces suggest (Hammond private communication, 1981):

\[
\frac{H_{m} - H_{a}}{H_{e} - H_{a}} = 4.09 \left( \frac{X_{sp}}{S_{i}} \right)^{-0.5}
\]  

(2.31)

\[
\frac{H - H_{a}}{H_{m} - H_{a}} = 1 - \frac{y}{\delta}
\]  

(2.32)

where

- \( H \) is the time-averaged enthalpy/temperature,
- \( H_{e} \) is the enthalpy/temperature at slot inlet,
- \( H_{a} \) is the ambient enthalpy/temperature and
- \( H_{m} \) is the maximum enthalpy/temperature.

2.6.1.2 Boundary condition for Xsp and outlet

Along the line Xsp and the room outlet plane a zero streamwise gradient was imposed to all the variables except \( U \) at Xsp and \( V \) at the outlet, which were specified as zero. Thus,
at the outlet: \( \partial \phi / \partial x = 0 \) and \( V = 0 \)
and at \( \text{Xsp} \): \( \partial \phi / \partial y = 0 \) and \( U = 0 \) 

(2.33)

2.6.2 At Walls, Floor and Ceiling:

In the present work the solid surfaces are stationary and impervious so the velocity components are specified zero. The enthalpy/temperature is therefore given a prescribed value. However, the kinetic energy and its dissipation rate requires special treatment as described in the next section.

2.7 WALL FUNCTION:

The simple boundary specification of zero velocity components at a stationary, non-slip and impermeable wall is valid, but with difficulties in numerical calculations when the flow is turbulent. Viscous and turbulent stresses are of the same order of magnitude near the wall. Thus the effective viscosity and other transport properties, fall to their laminar values and the result is a steep, non-linear variation with distance from the wall in velocities, scalar properties, turbulent viscosity and their gradients. It is still possible to compute these variations by packing the grid nodes into regions of steep gradient changes, close enough for accurate numerical calculation. However, this necessitates both a large computer and running costs. Also those functions appearing in turbulence model equations must properly represent the influences of the local Reynolds number of turbulence \( \mu_{eff}/\mu_\ell \), which for the type of turbulence model used in the present work is inappropriate. Fortunately, very close to the wall, one-dimensional “Couette flow” behaviour is observed, which offers a way around this dilemma in the form of a “bridging-over” across the near wall region, via the introduction of “wall-functions”. These functions are often expressed in the form of algebraic equations. It is therefore convenient to use a “wall-function” to connect the flux of \( \phi \) to the wall with the local \( \phi \)-difference across the Couette flow region and with other relevant quantities.

The wall function adopted for the present work is the same as the one Pun and Spalding (ref 31) have incorporated in their elliptic code, which is described in some detail by Launder and Spalding (ref 24).
a) **Momentum flux near the wall**

The velocity variation is assumed to conform to the following modified form of the logarithmic law:

\[ U^+ = \frac{U}{U_T} = \frac{1}{\kappa} \ln (E Y^+) \]  \hspace{1cm} (2.34)

where

- \( U \) is the velocity parallel to the wall at a distance \( y \) from the wall,
- \( U_T \) is the so-called "friction" velocity \( (\equiv \frac{\tau_w}{\rho}) = \frac{\tau_w}{\rho C_D^{\frac{1}{4}} K^\frac{1}{4}} \),
- \( \kappa \) & \( E \) are the law of the wall constants, whose values are 0.41 and 8.4 respectively (ref 33),
- \( Y^+ \) is the dimensionless distance from the wall \( (\equiv \frac{\rho U_T y/\mu_L}{C_D^{\frac{1}{4}} \rho K^\frac{1}{4} y/\mu_L}) \),
- \( \tau_w \) is the wall shear stress.

b) **Heat flux near the wall**

The distribution of enthalpy is assumed to be analogous to that for velocity given by equation 2.34. It may be written in the form:

\[ H^+ = \sigma_t (U^+ + P) \]  \hspace{1cm} (2.35)

where \( H^+ \) is defined in terms of the enthalpy near the wall \( (H) \), enthalpy at the wall \( (H_w) \), and the local flux to the wall \( (q_w) \) as follows:

\[ H^+ = \frac{(H - H_w) C_D^{\frac{1}{4}} K^\frac{1}{4}}{q_w} \]  \hspace{1cm} (2.36)

and \( P \) is the viscous sublayer "resistance". Jatatileke (ref 51) evaluated the \( P \)-function from experimental data for moderate to high molecular Prandtl numbers, and found that it is well represented by:
\[ p = 9.0 \left( \frac{\sigma_{\text{e}}}{\sigma_{\text{t}}} - 1 \right) \left( \frac{\sigma_{\text{e}}}{\sigma_{\text{t}}} \right)^{-1} \]  

(2.37)

where \( \sigma_{\text{e}} \) and \( \sigma_{\text{t}} \) are the laminar (molecular) and turbulent Prandtl numbers.

c) **Kinetic energy near the wall**

The kinetic energy is obtainable from its derivative, but with the diffusion of \( K \) to the wall set to zero, thus:

\[ \left( \frac{\partial K}{\partial y} \right)_{\text{wall}} = 0 \]  

(2.38)

The generation and dissipation terms are modified to be consistent with the known results for a Couette flow region. Details will be given in chapter 3.

d) **Dissipation rate near the wall**

Near the wall the length scale is assumed proportional to the distance from the wall, thus the dissipation rate can be expressed as:

\[ \varepsilon = \frac{C_{p} K_{y}^{3/2}}{K_{y}} \]  

(2.39)

using a variation on equation (2.30).

2.8 **THE GENERALIZED FORM OF THE EQUATIONS TO BE SOLVED:**

The equations to be solved can be expressed by a single equation of the form:

\[ \frac{\partial}{\partial x_{j}} (\rho \phi U_{j}) = \frac{\partial}{\partial x_{j}} \left( \Gamma_{\phi} \frac{\partial \phi}{\partial x_{j}} \right) + S_{\phi} \]  

Convection \hspace{.5cm} diffusion \hspace{.5cm} Source

terms \hspace{1cm} terms \hspace{1cm} terms
The expressions for diffusion coefficient, \( \Gamma_\phi \), and source term, \( S_\phi \), corresponding to a particular variable \( \phi \) are given in table 2.2, and the buoyancy production (ref 26,35,36,52-54) is given by:

\[
G_B = - \beta \, g \, \frac{\mu_t}{\sigma_t} \, \frac{\partial \theta}{\partial y}
\]  

(2.41)

The effective viscosity \( \mu_{\text{eff}} \) is defined by:

\[
\mu_{\text{eff}} = \mu_g + \mu_t
\]

(2.42)

and the effective exchange coefficient for H is given by:

\[
\Gamma_{H,\text{eff}} = \frac{\mu_g}{\sigma_g} + \frac{\mu_t}{\sigma_t}
\]

(2.43)

where

- \( \theta \) is the excess temperature \( (T - T_0) \),
- \( T_0 \) is some reference value,
- \( \beta \) is coefficient of thermal expansion and
- \( g \) is gravitational acceleration.
<table>
<thead>
<tr>
<th>Conserved property</th>
<th>$\phi$</th>
<th>$\Gamma_{\phi}$</th>
<th>$S_{\phi}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuity</td>
<td>1</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>x-direction momentum</td>
<td>U</td>
<td>$\mu_{\text{eff}}$ - $\frac{\partial P}{\partial x} + \frac{\partial}{\partial x} (\mu_{\text{eff}} \frac{\partial U}{\partial x}) + \frac{\partial}{\partial y} (\mu_{\text{eff}} \frac{\partial V}{\partial x})$</td>
<td></td>
</tr>
<tr>
<td>y-direction momentum</td>
<td>V</td>
<td>$\mu_{\text{eff}}$ - $\frac{\partial P}{\partial y} + \frac{\partial}{\partial x} (\mu_{\text{eff}} \frac{\partial U}{\partial y}) + \frac{\partial}{\partial y} (\mu_{\text{eff}} \frac{\partial V}{\partial y}) + \rho \beta g \theta$</td>
<td></td>
</tr>
<tr>
<td>Thermal energy</td>
<td>H</td>
<td>$\Gamma_{h,\text{eff}}$</td>
<td>$S_h$</td>
</tr>
<tr>
<td>Turbulence energy</td>
<td>K</td>
<td>$\mu_{\text{eff}}/\sigma_k$</td>
<td>$G_k - \rho \varepsilon + G_B$</td>
</tr>
<tr>
<td>Dissipation rate</td>
<td>$\varepsilon$</td>
<td>$\frac{\mu_{\text{eff}}}{\sigma_k}$</td>
<td>$\frac{\varepsilon}{K} ((G_k + G_B) C_1 - C_2 - \rho \varepsilon)$</td>
</tr>
</tbody>
</table>

Table 2.2: Exchange Coefficients and Source Terms
CHAPTER 3
NUMERICAL SOLUTION PROCEDURE FOR THE HIGHER-LEVEL METHOD

3.1 INTRODUCTION:

The "elliptic" governing partial differential equations obtained in Chapter 2 are complex and strongly inter-linked with no obvious equation available for the static pressure that appears in the momentum equations. An analytical solution is therefore not possible, and it is necessary to use numerical methods of solution. There are two main numerical solution techniques for the type of elliptic equations encountered in the present problem: "finite-difference" and "finite-element" methods. The main differences between these two methods concern the manner in which the discretized equations are derived. The benefit of finite-element methods (FEM) lies in its ability to fit irregular boundaries fairly readily, although difficulties have been experienced in bridging the steep property gradients encountered near walls in turbulent flows. There are a number of other technical difficulties in using the FEM for fluid flow (as opposed to solid mechanic) problems, and these are discussed by Patankar (ref 55). Fortunately, the flow domains considered here are all rectangular, and the finite-difference approach has been adopted.

3.2 FORMULATION OF THE FINITE-DIFFERENCE EQUATIONS:

3.2.1 Derivation Methods:

There are several ways to obtaining finite-difference equations (FDE's) from differential equations, some of the most common methods are briefly described below, and further details are given by Patankar (ref 55):

a) Taylor-series formulation:

In this method, the derivatives in the differential equations are obtained via a truncated Taylor series expansion. An assumption that the variation of \( \phi \) in a particular direction may be represented by polynomial is also invoked.

The method offers less flexibility and little physical insight into the derivation of the FDE's, and the resultant equations do not always satisfy the conservation requirement.
b) Variational formulation:

Method is based on the calculus of variations. It shows that, the solution of a certain differential equation is equivalent to minimizing a related quantity called "functional". This equivalence is known as variational principle. This method is complex and rather limited in applicability.

c) Weighted residuals:

This is a powerful method for solving the differential equations. The method is summarized below, although it is described by Patankar (ref 55) and in more detail by Finlayson (ref 56).

Supposing the differential equation is of the form:

$$L(\phi) = 0 \quad (3.1)$$

and

$$\phi = a_0 x^0 + a_1 x^1 + a_2 x^2 + \ldots + a_n x^n \quad (3.2)$$

where $a$'s are parameters. The substitution of equation (3.2) into the equation (3.1) gives a residual, which is defined as:

$$R = L(\phi) \quad (3.3)$$

This residual should be as small as possible. Let propose that

$$\int_{W} R \, dx = 0 \quad (3.4)$$

where $W$ is a weighting functions. A number of equations can be generated for evaluating the unknown parameters, by first integrating equation (3.4) over the domain of interest, and then choosing a succession of weighting functions. Different versions of the method result from the choice of weighting functions. A simplest weighting function is obtained when $W=1$. A number of weighting residual equations may be generated by dividing the complete solution domain into subdomains or control-volumes, and setting the weighting function to be unity over one control volume at a time and zero elsewhere. This version of weighted residuals is called the "subdomain", "micro-integration" or "control-volume" method. The advantage of this approach is that the approximations inherent in the discretization process are easily identifiable, and it is easy to ensure that the resulting
FDE's satisfy the conservation requirement. This method has been chosen for the present work.

3.2.2 Computational Grid and Control-Volume:

The first step in the derivation of the FDE's is the establishment of a discritization grid, and the definition of the points at which variables are located. The grid spacing may be non-uniform, to allow the spacing to be small in regions with steep property gradients, such as near walls. The intersection of the grid lines are grid nodes. Each grid node is assumed to be enclosed in its own cell or control volume. The control volume boundaries were taken to be mid-way between the grid nodes in the present study.

Fig. (3.1) illustrates the two-dimensional grid applied in the present work. A typical grid node P and its neighbours N, S, E and W are shown in fig. (3.2) as well as the relevant control volume (i.e., the volume enclosed by the dashed lined). The boundaries of the control volume are labelled with the lower-case version of the label of the neighbour nodes (n, s, e and w).

3.2.3 Location of Dependent Variables:

All dependent variables are stored at the grid nodes, except the velocity components which are calculated at "staggered" locations (i.e. mid-way between adjacent grid nodes), as shown in fig. (3.3). This "staggered" grid, which was first used by Harlow and Welch (ref 57) and subsequently adopted by Patankar and Spalding (ref 29) in their SIMPLE algorithm, has the advantage of ensuring that the velocities are directly available for calculating the convective fluxes of scalar flow variables as well as lying between the location of the static pressures which "drive" them. The staggered location and control volume for velocity components are shown in fig. (3.4) and (3.5).

3.2.4 The General Finite-Difference Equations:

The governing differential equations which must be replaced by FDE's are:

(a) Continuity

\[
\frac{\partial}{\partial x_j} (\rho U_j) = 0
\]
and,

(b) the general differential equation

\[ \frac{\partial}{\partial x_j} (\rho \phi u_j) = \frac{\partial}{\partial x_j} (\Gamma \phi \frac{\partial \phi}{\partial x_j}) + s_\phi \]

(3.6)

Both equation (3.5) and (3.6) may be written, for an incompressible flow, in the form:

\[ \rho u_j \left( \frac{\partial \phi}{\partial x_j} \right) = \frac{\partial}{\partial x_j} (\Gamma \phi \frac{\partial \phi}{\partial x_j}) + s_\phi \]

(3.7)

Equation (3.6) can alternatively be written in the form:

\[ \frac{\partial}{\partial x_j} (J_j) = s_\phi \]

(3.8)

where \( J_j \) is the total flux in direction \( j \), which defined as:

\[ J_j = \rho u_j \Gamma \phi \frac{\partial \phi}{\partial x_j} \]

(3.9)

The integration of equation (3.8) over the control volume shown in Fig (3.2) gives:

\[ J_e - J_w + J_n - J_s = s_\phi \Delta x \Delta y \]

(3.10)

where \( J_e, J_w, J_n \) and \( J_s \) are the integrated total fluxes over the control-volume faces. The source term, \( s_\phi \), may simply be written in quasi-linear form as:
\[ S_\phi = S_p + S_p' \phi_p \]  
\[ (3.11) \]

where \( S_p \) and \( S_p' \) are determined from a suitable finite difference approximation of the \( S_\phi \) expression for the particular variable \( \phi \). Hence, equation (3.10) becomes:

\[ J_e - J_w + J_n - J_s = (S_p + S_p' \phi_p) \Delta x \Delta y \]  
\[ (3.12) \]

The continuity equation (3.5) may also be integrated over the control volume, and the result can be written in the form:

\[ F_e - F_w + F_n - F_s = 0 \]  
\[ (3.13) \]

where \( F_e \), \( F_w \), \( F_n \) and \( F_s \) are the mass flow rate through the control volume faces:

\[ F_e = (\rho U)_e \Delta y \]
\[ F_w = (\rho U)_w \Delta y \]
\[ F_n = (\rho V)_n \Delta x \]
\[ F_s = (\rho V)_s \Delta x \]  
\[ (3.14) \]

By multiplying the equation (3.13) by \( \phi_p \) and subtracting from equation (3.12), the following equation will result:

\[ (J_e - F_e \phi_p) - (J_w - F_w \phi_p) + (J_n - F_n \phi_p) - (J_s - F_s \phi_p) = (S_p + S_p' \phi_p) \Delta x \Delta y \]  
\[ (3.15) \]

This is in fact the finite-difference equivalent to equation (3.7). According to Patankar (ref 55), terms \( (J_e - F_e \phi_p) \), etc... can be replaced by:
After substituting the relations (3.16) into equation (3.15) and some rearrangement, the FDE has the final form:

\[
\sum_{E,W,N,S} a_{\phi} \phi_{P} = \left( \sum_{E,W,N,S} a_{\phi} \right) + b
\]

(3.17)

where

\[
a_{E} = D_{e} L \left( |P_{e}| \right)_{e} + \text{Max} \left( -F_{e}, 0 \right)
\]

\[
a_{W} = D_{w} L \left( |P_{w}| \right)_{w} + \text{Max} \left( F_{w}, 0 \right)
\]

\[
a_{N} = D_{n} L \left( |P_{n}| \right)_{n} + \text{Max} \left( -F_{n}, 0 \right)
\]

\[
a_{S} = D_{s} L \left( |P_{s}| \right)_{s} + \text{Max} \left( F_{s}, 0 \right)
\]

\[
a_{p} = \left( \sum_{E,W,N,S} a_{\phi} \right) - S_{p} \Delta x \Delta y
\]

\[
b = S_{p} \Delta x \Delta y
\]

(3.18)

The \( D_{e} \), \( D_{w} \), \( D_{n} \) and \( D_{s} \) are diffusion terms and are defined as:

\[
D_{e} = \Gamma_{\phi,e} \Delta y / (\delta x)_{e}
\]

\[
D_{w} = \Gamma_{\phi,w} \Delta y / (\delta x)_{w}
\]

\[
D_{n} = \Gamma_{\phi,n} \Delta x / (\delta y)_{n}
\]

\[
D_{s} = \Gamma_{\phi,s} \Delta x / (\delta y)_{s}
\]

(3.19)
Peₑ, Peᵦ, etc... are Peclet numbers representing the ratio of the strengths of the convection and diffusion terms:

\[ Peₑ = (\rho U \delta x / \Gamma_\phi)_ₑ = \frac{Fₑ}{Dₑ} \]
\[ Peᵦ = (\rho U \delta x / \Gamma_\phi)_ᵦ = \frac{Fᵦ}{Dᵦ} \]
\[ Peₙ = (\rho V \delta y / \Gamma_\phi)_ₙ = \frac{Fₙ}{Dₙ} \]
\[ Peₛ = (\rho V \delta y / \Gamma_\phi)_ₛ = \frac{Fₛ}{Dₛ} \]  

(3.20)

The exchange coefficients \( \Gamma_\phi \) at control-volume faces are simply specified as:

\[ \Gammaₚₑ = (\Gammaₚₑ + \Gammaₑₑ) / 2 \]
\[ \Gammaₚᵦ = (\Gammaₚᵦ + \Gammaₑᵦ) / 2 \]
\[ \Gammaₚₙ = (\Gammaₚₙ + \Gammaₑₙ) / 2 \]
\[ \Gammaₚₛ = (\Gammaₚₛ + \Gammaₑₛ) / 2 \]  

(3.21)

L(1Pe₁) is a function related to the particular differencing scheme adopted for the finite-difference procedure. The \( \text{Max}(A,B) \) denotes the greater of A and B.

3.2.5 The Differencing Schemes:

In order to illustrate the choice of differencing schemes, a steady one-dimensional situation in which only the convection and diffusion terms exist will be considered. The governing differential equation then simplifies to:

\[ \frac{d}{dx} (\rho U \phi) = \frac{d}{dx} \left( \Gamma_\phi \frac{d\phi}{dx} \right) \]  

(3.22)
and the continuity equation to:

\[ \frac{d}{dx} (\rho U) = 0 \quad \text{or} \quad \rho U = \text{const.} \]  

(3.23)

The corresponding grid nodes are illustrated in figure (3.6). Finally, the integration of equation (3.22) over the control volume in figure 3.6 yields:

\[ (\rho U\phi)_e - (\rho U\phi)_w = \left[ \Gamma_{\phi} \frac{\partial \phi}{\partial x} \right]_e - \left[ \Gamma_{\phi} \frac{\partial \phi}{\partial x} \right]_w \]  

(3.24)

a) Central Difference Scheme

In this scheme a piecewise-linear variation for \( \phi \) is assumed between the grid nodes, therefore the value of \( \phi \) at an interface is specified as:

\[ \phi_e = (\phi_E + \phi_P)/2 \]

\[ \phi_w = (\phi_P + \phi_W)/2 \]  

(3.25)

and the diffusion terms are given in the form:

\[ \left( \Gamma_{\phi} \frac{d\phi}{dx} \right)_e = \Gamma_{\phi,e} \frac{\phi_E - \phi_P}{(\delta x)_e} \]

\[ \left( \Gamma_{\phi} \frac{d\phi}{dx} \right)_w = \Gamma_{\phi,w} \frac{\phi_P - \phi_W}{(\delta x)_w} \]  

(3.26)

Thus, equation 3.24 may be written in the form:

\[ \frac{1}{2}(\rho U)_e (\phi_E + \phi_P) - \frac{1}{2}(\rho U)_w (\phi_P + \phi_W) = \]

\[ \Gamma_{\phi,e} (\phi_E - \phi_P)/(\delta x)_e - \Gamma_{\phi,w} (\phi_P - \phi_W)/(\delta x)_w \]  

(3.27)
After some rearrangement, and using equation 3.23, the finite-difference form of equation 3.22 becomes:

\[ a_{p} \phi_{p} = a_{E} \phi_{E} + a_{W} \phi_{W} \]  

(3.28)

where

\[ a_{E} = D_{e} - \frac{F_{e}}{2} \]
\[ a_{W} = D_{w} + \frac{F_{w}}{2} \]
\[ a_{p} = a_{E} + a_{W} \]  

(3.29)

According to Patankar (ref 55), Runchal (ref 58), and others, for given values of \( h \) and \( \beta \) the central difference scheme yields accurate results only when \( P_{e} < 2 \). Outside this limit the scheme may become inaccurate and, with an iterative method of solution, instability may ensue (ref 22,55,58,59).

b) Upwind Scheme

In this scheme the formulation of diffusion term is the same as with central difference scheme, except that the convection term is calculated in a different way (ref 55,60). The value of \( \phi \) at an interface is assumed to be equal to the value of \( \phi \) at the grid node on the "upstream" or "upwind" side of the face: for example

\[ \phi_{e} = \phi_{p} \quad \text{if} \quad F_{e} > 0 \]
\[ \phi_{e} = \phi_{E} \quad \text{if} \quad F_{e} < 0 \]  

(3.30)

When equations (3.25) is replaced by this definition, the \( a \)'s coefficients in equation 3.28 become:

\[ a_{E} = D_{e} + \text{Max} (-F_{e},0) \]
\[ a_{W} = D_{w} + \text{Max} (F_{w},0) \]  

(3.31)
The upwind scheme gives realistic and stable behaviour at all Peclet numbers. It is also superior to central difference scheme in terms of accuracy for $P_e > 2$, but slightly inferior for smaller values (ref 58).

c) Hybrid Scheme

This is a combination of central difference and upwind schemes. It employs the central difference scheme for $-2 < P_e < 2$, and the upwind scheme, in which the diffusion has been set equal to zero for outside this range. For example

$$\frac{a_E}{D_e} = -P_e \quad \text{if } P_e < -2$$

$$\frac{a_E}{D_e} = 1 - \frac{P_e}{2} \quad \text{if } -2 < P_e < 2$$

$$\frac{a_E}{D_e} = 0 \quad \text{if } P_e > 2$$

(3.32)

In combined form, the coefficient $a_E$ and $a_W$ have the form:

$$a_E = \max \left( -F_e, D_e - \frac{F_e}{2} \right)$$

$$a_W = \max \left( F_w, D_w + \frac{F_e}{2} \right)$$

(3.33)

d) Power-Law Scheme

According to Patankar (ref 55,61) the departure of the hybrid scheme from the exact result is large at $P_e = \pm 2$ (see fig 3.7). He felt that the setting of the diffusion terms to zero as soon as $|P_e|$ exceeds 2 as rather premature, and therefore proposed a power-law scheme which may be written in the form:

$$\frac{a_E}{D_e} = -P_e \quad \text{if } P_e < -10$$

$$\frac{a_E}{D_e} = (1 + 0.1 P_e)^5 - P_e \quad \text{if } -10 \leq P_e < 0$$

$$\frac{a_E}{D_e} = (1 - 0.1 P_e)^5 \quad \text{if } 0 \leq P_e \leq 10$$

$$\frac{a_E}{D_e} = 0 \quad \text{if } P_e > 10$$

(3.34)
In a combined form thus may be written as:

\[
\begin{align*}
\alpha_E &= D_e \max\{0, (1 - \frac{0.1 |F_e|}{D_e})^5\} + \max(0, -F_e) \\
\alpha_W &= D_w \max\{0, (1 - \frac{0.1 |F_w|}{D_w})^5\} + \max(0, F_w)
\end{align*}
\]  

(3.35)

It is now possible to specify the function \(L(|\text{Pe}|)\), which appears in equations 3.18, from the various schemes outlined above. These are summarised in table 3.1 below.

| Scheme         | \(L(|\text{Pe}|)\)                                      |
|----------------|--------------------------------------------------------|
| Central difference | 1 - 0.5 |\text{Pe}| |
| Upwind          | 1                                    |
| Hybrid          | \(\max(0, 1-0.5|\text{Pe}|)^5\)                        |
| Power law       | \(\max(0, (1-0.1|\text{Pe}|)^5\)                      |

All schemes discussed above give a physically realistic solution except the central differencing scheme, which requires a very fine grid in order to yield reasonable solution for \(\text{Pe} < 2\). The upwind scheme is adopted in the present work, as it appears to exhibit a reasonable balance between accuracy and economy. However, the other differencing schemes have been incorporated into the program structure, and could readily be used if greater numerical accuracy is required.

3.2.6 Special Attention On Momentum Equations:

The momentum equations can be obtained from equation 3.17 for the general variable \(\phi\), where \(\phi\) now stands for the relevant velocity components and \(\Gamma_\phi\) and \(S_\phi\) are given in table 2.2. The only complication arises due to the use of staggered grid and control volume (see fig. 3.3-3.5) for U and V components. The use of staggered grid allows the pressure differences \((P_P-P_E \text{ & } P_P-P_N\) ) to be employed directly to calculate the pressure force acting
on the control volume for each velocity component. The resulting U-momentum and V-momentum equations have the form:

\[ a_p U_p = \left( \sum_{E, W, N, S} aU \right) + b + A_e (P_p - P_E) \]

\[ a_p V_p = \left( \sum_{E, W, N, S} aV \right) + b + A_n (P_p - P_N) \]

(3.36)

The source term \(b\) is defined in the same manner as in equations 3.18, where the pressure gradient is not included in the source term quantities \((S_p \& S_P)\). The pressure gradients \(A_e(P_p - P_E)\) & \(A_n(P_p - P_N)\) are acting in the U and V control volume respectively, where:

\[ A_e = \Delta y \times 1 \quad , \quad A_n = \Delta x \times 1 \]

(3.37)

The momentum equations cannot be solved unless the pressure field is given or estimated, and the resulting velocities will not satisfy the continuity equation unless they are based on the correct pressure field. However, 'imperfect' velocity components may be estimated from guessed pressures, \(P^*\), from the solution of the following equations:

\[ a_p U^* = \left( \sum_{E, W, N, S} aU^* \right) + b + A_e (P^*_p - P^*_E) \]

\[ a_p V^* = \left( \sum_{E, W, N, S} aV^* \right) + b + A_n (P^*_p - P^*_N) \]

(3.38)

where \(U^*\) and \(V^*\) are called 'starred' velocity components.

3.2.7 The Pressure and Velocity Corrections:

The guessed or starred pressure \(P^*\) must be improved in such a way that more accurate starred velocity components \((U^* \& V^*)\) are produced, and therefore get closer to satisfying the continuity equation. Correction quantities such as \(P^*, U^*\) and \(V^*\) whose purpose is to bring the continuity and momentum equation into balance, may be determined as follows:
\[ P = P* + P' \]
\[ U = U* + U' \]
\[ V = V* + V' \]  \hspace{1cm} (3.39)

The substitution of equations 3.39 into equations 3.38, and the substraction from 3.36, leads to:

\[ a_U U' = (\sum_{E,W,N,S} aU') + A_e (P'_p - P'_e) \]
\[ a_P V' = (\sum_{E,W,N,S} aV') + A_n (P'_p - P'_N) \]  \hspace{1cm} (3.40)

In the SDTLE algorithm of Patankar and Spalding (ref 29) adopted here, the terms $\sum a U'$ and $\sum a V'$ are neglected from the above equations, for the reason given in section 3.2.8. Equation 3.40 then becomes:

\[ a_P U'_p = A_e (P'_p - P'_e) \]
\[ a_P V'_p = A_n (P'_p - P'_N) \]  \hspace{1cm} (3.41)

or

\[ U'_p = d_e (P'_p - P'_e) \]
\[ V'_p = d_n (P'_p - P'_N) \]  \hspace{1cm} (3.42)

where

\[ d_e = A_e/a_P \quad , \quad d_n = A_n/a_P \]  \hspace{1cm} (3.43)

The substitution of equations 3.42 into equations 3.39 yield:

\[ U_p = U* + d_e (P'_p - P'_e) \]
\[ V_p = V* + d_n (P'_p - P'_N) \]  \hspace{1cm} (3.44)
3.2.8 The Pressure-Correction Equation

The substitution of equations 3.44 into the continuity equation (3.13), and rearranging yields the desired equation for the pressure-correction \( P^* \):

\[
ap^p = \left( \sum_{E,W,N,S} a_i \right) + b
\]

where

\[
a_E = \rho_e \frac{\partial}{\partial y} A_e
\]
\[
a_W = \rho_w \frac{\partial}{\partial x} A_w
\]
\[
a_N = \rho_n \frac{\partial}{\partial n} A_n
\]
\[
a_S = \rho_s \frac{\partial}{\partial s} A_s
\]
\[
a_P = a_E + a_W + a_N + a_S
\]
\[
b = (\rho U^* A)_w - (\rho U^* A)_e + (\rho V^* A)_s - (\rho V^* A)_n
\]

(3.45)

(3.46)

Where

\[A_e = A_w = \Delta y\]
\[A_n = A_s = \Delta x\]

(3.47)

and \( \rho_e \), \( \rho_w \), \( \rho_n \) and \( \rho_s \) can be calculated by interpolation from the value of the densities at the grid nodes.

The source term in equation 3.46, \( b \), represents a mass balance on the control volume based on the preliminary velocities in the calculation domain. The purpose of pressure correction is to reduce this "mass source" to zero. Since the mass-source and pressure corrections will become zero after the velocities and the pressure satisfy the momentum and continuity equations, the final solution will be independent of the terms \( \sum a_U' \) and \( \sum a_V' \) neglected from equations 3.40.
Equation 3.45 has the same form as 3.17 and can be solved in a similar manner. The resulting pressure-correction may then be used to adjust the velocity components from equations 3.44 and pressure from equation 3.39.

3.3 THE INSERTION OF BOUNDARY CONDITIONS:

The FDE's are not applicable for boundary nodes since they refer to the points which lying outside the calculation domain. The control volumes for different variables near to boundary are shown in fig 3.8. This arrangement has the advantage for meeting the conservation requirement and facilitating the flux calculation.

A particular expression for $\phi_N$ (see fig 3.9), which coincides with the face n of the adjoining control volume can be inserted to break the normal link between $\phi_P$ and $\phi_N$ in the FDE's. This link breaks simply by specifying the appropriate coefficient in FDE to zero (for example $a_N=0$).

The boundary appropriate to the present study were outlined in Section 2.6. The special practice which are required to incorporate the appropriate boundary conditions for the various flow variables in the wall region will be outlined in this Section.

3.3.1 Tangential Velocity:

The tangential velocity for the node next to the wall (for example, $U_p$ in fig 3.9b) can be obtained from the usual FDE for momentum. However, $U_p$ follows the logarithmic law of the wall (equation 2.34), which relates it to friction velocity $U_T$. It is necessary to incorporate the correct shear stress into the FDE. The shear stress may be introduced into the FDE by breaking the link between the node on the wall and the adjacent grid node, and by modifying the source terms in FDE. For example, for nodes P and N shown in fig 3.9b

$$a_N = 0$$

If point P is within the turbulent region ($Y^+ > 11.5$), the correct shear stress can then be determined via the logarithmic law, equation 2.34:
\[ \tau_w = \kappa C_D^{1/4} \rho U_p K_P^{1/2} \ln \left( E \rho \delta K^{1/2} C_D^{1/4} / \mu \right) \]  

(3.49) 

where 

\[ Y^+ = C_D^{1/4} \rho K_P^{1/2} y / \mu \]  

(3.50) 

However, if point \( P \) is within the viscous sublayer \((Y^+ < 11.5)\), then the shear stress may be simply evaluated from: 

\[ \tau_w = \mu \frac{U_p}{\delta} \]  

(3.51) 

The appropriate form of \( \tau_w \) obtained from either of these expressions can then be inserted into the FDE via the source terms as follows (ref 31): 

\[ S_P = 0 \] 

\[ S_P' = - \tau_w A_n / U_P \]  

(3.52) 

It should be noted that no special treatment is required for the velocity normal to the wall. 

3.3.2 Heat Flux Near The Wall: 

The enthalpy boundary condition can be introduced to the thermal energy FDE in a similar manner to that for velocity. Equation 2.36 can also be written in terms of the wall shear stress as follows (ref 31): 

\[ St \equiv \frac{-q_w}{(H_P - H_N) \rho |U_p|} = \frac{\tau_w / (\rho U_P^2)}{\sigma_t (1 + \rho \sqrt{\tau_w / (\rho U_P^2)}} \]  

(3.53)
where, if the node $P$ lies within the turbulent region, $\tau_w$ is evaluated from equation 3.49, or, if it is within the viscous sublayer, from equation 3.51. The FDE coefficient $a_n$ is first set to zero (see fig 3.9a), and the heat flux is then incorporated into the FDE through the source terms as before (ref 31):

$$S_p = \left| St \rho U_p A_n \right| H_N$$

$$S'_p = \left| St \rho U_p A_n \right|$$

### 3.3.3 Turbulence Kinetic Energy

To achieve the conditions which are mentioned in Section 2.7(c), it is necessary again to set $a_n$ to zero (see fig 3.9a). Due to the steep velocity gradients near the walls, the source terms are best expressed through shear stress (ref 31). These terms may therefore be written in the form:

$$S_k = G_k - \rho \varepsilon$$

$$= \tau_w \frac{\partial U}{\partial y} - \frac{C_D \rho^2 K^2}{\mu_t}$$

$$= \frac{\tau_w \partial U}{\partial y} - \frac{C_D \rho^2 K^2}{\tau_w} \frac{\partial U}{\partial y}$$

The integration of $S_k$ over the control volume shown in fig 3.9a yields:

$$\int \int S_k dV = \tau_w U_s A_n - C_D \rho^2 K_p U_s A_n / \tau_w$$

hence,

$$S_p = \tau_w U_s A_n$$

$$S'_p = C_D \rho^2 K_p A_n / \tau_w$$

(3.54)
3.3.4 The Dissipation Rate:

The dissipation rate \( \varepsilon_P \) (see fig 3.9a) is obtained directly from the expression 2.39, thus:

\[
\varepsilon_P = \frac{C_D^{3/4} K_P^{3/2}}{\kappa \delta}
\]

(3.56)

The value of \( \varepsilon_P \) is incorporated into the FDE for the near-wall node through the source terms (ref 31,55):

\[
S_p = \alpha \varepsilon_P
\]

\[
S_p' = -\alpha
\]

(3.57)

where \( \alpha \) is a very large number (eq 10).

3.4 THE SOLUTION OF THE FINITE-DIFFERENCE EQUATIONS:

Equations 3.17 and 3.36 must be solved to obtain \( U, V, K, \varepsilon, H \) and \( P' \) for all the grid nodes in the calculation domain. These equations must be solved simultaneously and, since they are non-linear and inter-linked, it is necessary to employ an iterative solution procedure. The particular iterative method adopted in the present work is a combination of "line-by-line" iteration with a "block" line adjustment applied to each line. For each grid node on a line considered (see fig 3.10) the FDE's can be expressed as:

\[
\phi_j = A_j \phi_{j+1} + B_j \phi_{j-1} + C_j
\]

(3.58)

where \( j \) denotes the position along a particular line, in the range from \( j=2 \) to \( j=n-1 \). The boundary conditions (i.e., node positions \( j=1 \) and \( j=n \)) are incorporated in \( C_j \). With respect to equation 3.17:

\[
\phi_j = \phi_P \quad \phi_{j+1} = \phi_N \quad \phi_{j-1} = \phi_S
\]
Equations 3.58 for nodes on a line are solved simultaneously by matrix-inversion procedure. While the $\phi$ values on a line are being solved, the $\phi$ values on both sides of the line are kept unchanged. The computer code employed in the present study has been developed from that of Pun and Spalding (ref 31), who adopted the Tri-Diagonal Matrix Algorithm (TDMA), based on an elimination procedure derived by successive substitution (ref 62) for solving elliptic FDE sets.

An alternative way of rewriting equation 3.58, obtained by elimination, is:

$$\phi_j = A''_j \phi_{j+1} + B''_j$$

$$j = n-1, n-2, \ldots, 2$$

(3.60)

and the transformed coefficients $A''_j$ & $B''_j$ are given by the recurrence relations:

$$A''_j = A_j / (1 - A''_{j-1} B_j)$$

$$B''_j = (C_j + B''_{j-1} B_j) / (1 - A''_{j-1} B_j)$$

(3.61)

3.4.1 The Block-adjustment Procedure:

The block-adjustment procedure is used in an effort to successfully obtain a converged solution. The idea is to add a uniform increment $\delta \phi$ to the values of variables $\phi$ in each line, although the increment is varied from line to line. This
increment is applied in such a way that, after adjustment, all the lines satisfy the overall conservation requirement. Thus equation 3.17 can be written as:

\[ A_{ij}^P \phi_{ij} = A_{ij}^E \phi_{i+1,j} + A_{ij}^W \phi_{i-1,j} + A_{ij}^N \phi_{i,j+1} + A_{ij}^S \phi_{i,j-1} + B_{ij} \]

(3.62)

where the subscripts i and j denote the position in the x and y directions respectively (see fig 3.10).

If the prevailing solution procedure by line iteration is denoted by \( \phi_{ij} \), then a uniform increment \( \delta \phi_i \) is added to the interior nodes on each line, changing the equation 3.62 as follows:

\[ \sum_j A_{ij}^P (\phi_{ij} + \delta \phi_i) = \sum_j \left( A_{ij}^E \phi_{i+1,j} + A_{ij}^W \phi_{i-1,j} + A_{ij}^N \phi_{i,j+1} + A_{ij}^S \phi_{i,j-1} + B_{ij} \right) \]

(3.63)

Further manipulation of equation 3.63 results in the following equation for the \( \delta \phi_i \)'s:

\[ d_i \delta \phi_i = a_i \delta \phi_{i+1} + b_i \delta \phi_{i-1} + c_i \]

(3.64)

where

\[ a_i \equiv \sum_j A_{ij}^E, \quad b_i \equiv \sum_j A_{ij}^W \]
\[ C_i = \sum_j \left( A_{ij}^P \tilde{\phi}_{i,j} - A_{ij}^E \tilde{\phi}_{i+1,j} - A_{ij}^W \tilde{\phi}_{i-1,j} \\
- A_{ij}^N \tilde{\phi}_{i,j+1} - A_{ij}^S \tilde{\phi}_{i,j-1} - B_{ij} \right) \]

\[ d_i = \sum_j \left( A_{ij}^P - A_{ij}^N - A_{ij}^S \right) \]  

(3.65)

The equation 3.64 can be solved by TDMA with the increments \( \delta \phi \)'s set to zero at boundaries. The coefficient \( C_i \) is in fact the sum of residual sources from equation 3.17 for a line and, when \( C_j \) is zero, \( \delta \phi \) will be zero.

### 3.4.2 The Algorithm and the Sequence of Solution:

The solution procedure adopted for the present work is similar to the SIMPLE algorithm of Patankar and Spalding (ref 29), and more recently described by Patankar (ref 32), except that:

i) all variables are solved on a line before advancing the sweep, as advocated by Pun and Spalding (ref 31), and

ii) a block-adjustment procedure is employed to ensure continuity.

The sequence in which the FDE’s are solved is as follows:

a) Variables are initialized.

b) A line is considered.

c) Exchange coefficients \( \Gamma \) are defined (via equation 3.21).

d) Convection and diffusion terms are calculated (from equations 3.14 and 3.19).

e) The finite-difference coefficients are evaluated (from equation 3.18).

f) The FDE’s (equations 3.17 and 3.36) are solved by TDMA.

g) Block-adjustment coefficients are calculated (equation 3.65).

h) The steps b-g are repeated for all the variables except \( P \).

i) \( u^* \) and \( v^* \) are used to calculate the coefficients in pressure-correction equation (equation 3.46).

j) Pressure-correction equation (equation 3.45) is solved using the TDMA.
k) The resulting pressure corrections are used to update the pressure and velocity components in the manner indicated by equations 3.39 and 3.44.

l) The cycle of steps b-k represents one iteration. At the end of each iteration the equation for block-adjustment (equation 3.64) is solved by TDMA.

m) The increments \( \delta \phi \) are added to the corresponding \( \phi \)'s on each line.

h) Steps b-m are repeated until a converged solution is achieved.

3.5 NUMERICAL CONSIDERATIONS:

3.5.1 Convergence:

Equation 3.17 may also be written in the form:

\[
\phi_P = \frac{\sum_{E,W,N,S} a \phi}{a_P} + \frac{b}{a_P}
\]

(3.66)

According to Scarborough (ref 63), as discussed by Gosman et al (ref 22) and Patankar (ref 55), a successive substitution method ensures the convergence of a linear equation of a type such as equation 3.66 (i.e. when \( a \)'s and \( b \) are constants) if:

\[
\frac{\sum |a|}{E,W,N,S} \leq 1 \text{ for all equations}
\]

\[
\frac{|a_P|}{a_P} \leq 1 \text{ for at least one equation}
\]

(3.67)

All equations to be solved must satisfy this condition, thus:

\[
a_i > 0
\]

\[
a_P > \sum_{E,W,N,S} a
\]

(3.68)
However, in the present work the coefficients are not constant, and indeed they vary from one iteration to another. Fortunately Scarborough's criterion (Eq 3.67) is known to be a sufficient rather than a necessary condition for convergence and, when the above criteria are obeyed, it may be possible to solve non-linear equations. In an iterative method of solution, it is necessary to examine how the current solution approximates to the exact solution of the FDE's at the end of each iteration, for each dependent variable. The convergence criteria applied in the present work is based on the residual-sources of the FDE's. The residual source for equation 3.17, is defined by:

$$ R_\phi = a_\phi P - \sum_{E,W,N,S} a_\phi + b $$

(3.69)

For an exact solution, $R_\phi$ must be zero to satisfy the FDE, but for an inexact or nearly exact solution the normalized residual-source can be defined as:

$$ \sum_{\text{all field}} R_\phi \leq \lambda $$

$R_\phi, \text{ref}$

(3.70)

where $\lambda$ is a constant. $R_\phi, \text{ref}$ is a reference residual-source that is specified relative to an appropriate property at entrance of the flow. In the present study, the value for the convergence criteria $\lambda$ was typically set at less than 0.005.

The residual-source for pressure correction equation (3.45) was defined as the mass balance on the relevant control volume (i.e. $b$ in equation 3.46).

3.5.2 Stability:

The equations to be solved are non-linear and strongly interlinked, and must be solved simultaneously. They are therefore prone to numerical instability. Since iterative solution of linear and non-linear equations using the Scarborough criteria will normally converge, the instability must be due to the interlinkage between the dependent variables. The result of numerical instability is the non-satisfaction of the mass conservation on a local and overall basis. This leads to an oscillation of the residual-sources rather than convergence. Such oscillations are particularly prevalent in the case of the buoyant flow.
The major sources for instability identified in the present study were:

- non-linearity of the equations,
- velocity-pressure interlinkage of the momentum equations,
- buoyancy-linkage of momentum and thermal energy equations,
- buoyancy-linkage of $K$, $\varepsilon$ and thermal energy equations,
- buoyancy-linkage of $\mu_t$ through the $K$ and $\varepsilon$ equations.

The following techniques may reduce the numerical instability, although they proved only of rather limited success in buoyant flows.

(a) **Under-relaxation**

The use of some form of under-relaxation is very useful when solving non-linear equations. It reduces the magnitude of current error level in the field, while preventing the divergence in the iterative solution of equations. The way in which the dependent variables were under-relaxed in the present study is as follows:

Equation 3.17 may be written in the form:

$$
\phi_p = \sum_{E,W,N,S} \frac{a\phi}{a_p} b
$$

(3.71)

Supposing $\phi_p^*$ is the value of $\phi_p$ from the previous iteration, then by subtracting from both sides of equation 3.71, it becomes:

$$
\phi_p - \phi_p^* = \sum_{E,W,N,S} \frac{a\phi}{a_p} b - \phi_p^*
$$

(3.72)

Thus the right-hand side of the above equation gives the change in the variable between the current and previous iteration. This change can be modified by introducing a relaxation factor, $\xi$, as follows:
By rearrangement of the terms, equation 3.73 becomes:

\[ \frac{a_p}{\xi} \phi_p = \sum_{E,W,N,S} a_p \phi + b + (1 - \xi) \frac{a_p}{\xi} \phi^* \]

(3.74)

where

\[ 0 < \xi < 1 \] implies under-relaxation, and
\[ \xi = 1 \] means no relaxation.

The value of the relaxation factors for the different variables adopted in the present work are similar to those which were used by Pun and Spalding (ref 31) for the computation of turbulent flow through a sudden enlargement in a circular pipe. These relation factors are listed in table 3.2.

<table>
<thead>
<tr>
<th>Variable</th>
<th>U</th>
<th>V</th>
<th>K</th>
<th>( \varepsilon )</th>
<th>H</th>
<th>P</th>
<th>( \mu )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Under-relaxation</td>
<td>0.3</td>
<td>0.3</td>
<td>0.5</td>
<td>0.5</td>
<td>1.0</td>
<td>0.5</td>
<td>0.5</td>
</tr>
</tbody>
</table>

(b) Linearization:

The FDE source terms may induce the divergence of the iteration solution, if they are not properly linearized. The general approach to the linearization of the source terms was indicated by equation 3.11. The quantity \( S_p \) must be either zero or negative. Thus in the \( K \) & \( \varepsilon \) equations, the negative source terms are incorporated into \( S_p \). If some part of the source terms may change sign (such as the buoyancy source terms in the \( V \), \( K \) & \( \varepsilon \) equations) a suitable arrangement would be;
\[ S_p \approx \text{Max} (S_{\text{TERM}}, 0.) + \ldots \]
\[ S_p \approx -\text{Max} (-S_{\text{TERM}}, 0.) - \ldots \]

(3.75)

where \( S_{\text{TERM}} \) can be negative or positive.

(c) Line-traverse:

Increasing the number of TDMA traverses on a line might possibly yield a more complete solution of the equations during each iteration. In the present study, the maximum number of traverses on a line for all variables specified was two. However, if the largest of the absolute value of the algebraic sums of the residual sources for any variable on the line is smaller than a criteria value (say, 0.001) no more TDMA traverse were made on that line.

(d) Initialization:

Specifying realistic initial field values for all variables can be an important factor in ensuring both fast convergence and stability. In the present study the wall-jet profile analysis of Hammond (ref 33, 34) has been used to initialize the fields values for all variables (see fig 3.11) prior to the first iteration of the initial run. The converged solution for a 21 X 21 grid with an expansion between adjacent nodes of 1.1 on both the \( x \) and \( y \) directions is obtained after approximately 300 iterations, in the non-buoyant case. However, for subsequent runs, the results of previous computations can be used to initialize the field values for all variables, and this procedure reduces the number of iterations needed to obtain a converged solution to 30-200. The exact number of iterations depends on the degree of change in the input data from the previous run.

(e) Grid adjustment:

Generally the number of the grid lines must be sufficient to ensure numerical accuracy. In the case of a non-uniform grid, special attention must be paid to the ratio between two adjacent grid lines and the ratio between the cell walls. A series of tests for grid-dependence is therefore necessary, including a test of the influence of the number of grid nodes and one of the expansion ratio between neighbouring grid lines. These tests were conducted using various grids having between 225 and 729 nodes, and expansion ratios of 1.1 to 1.3 in both \( x \) and \( y \) directions. The results of these tests are presented in figures 3.12 to 3.14, where for the computations shown in figures 3.12 and 3.13 the expansion ratio between the grid lines was held at 1.1. It is clear that, although coarse 15 X 15 and 17 X 17 grids led to a converged solution, they differ from the results for the other, more refined grids. These differences are particularly significant in regard to the convective heat transfer coefficient (fig 3.12), but much less so in regard to the velocity field.
(fig 3.13). Apart from the coarse 15 X 15 and 17 X 17 grids, the results obtained using the various meshes are quite similar for most the variables in the field.

All the subsequent computed results presented in this thesis were obtained using a 21 X 21 finite-difference grid, as this gave rise to reasonable balance between rapid convergence and numerical accuracy when an expansion ratio of 1.1 was employed (see fig 3.14). A converged solution was typically achieved after 300 iterations for non-buoyant flows using a guessed initialization of the field variables. This corresponds to a central processor time of about 3300 seconds on the Cranfield GEC 4085 mini-computer.

3.5.3 Accuracy:

The accuracy of the solution obtained with the present method depends on the following factors:

i- The degree to which the solution satisfies the PDE's: this can be assessed by the level of the residual-sources defined by equation 3.69.

ii- The degree to which the solution depends on the grid arrangement: a grid independence test must be sought by increasing the number of grid lines, or concentrating them into regions where large variations in the variables occur, such that no significant change is observed with further increase.

iii- The conditions imposed at boundaries; the realistic adjustment of the variables at boundaries improves the degree of accuracy in the prediction.

iv- The adequacy of the turbulence model: the correctness of the turbulence model is open to question, and that may have a large influence to the accuracy of the problem.

Attempts have been made to reduce to a certain extent the errors due to factors i, ii and iii which are computation factors in origin, but no modifications have been made to the popular, fe turbulence model.

3.6 THE FLOW DIAGRAM FOR THE HIGHER-LEVEL COMPUTER PROGRAM:

Figure 3.15 shows the flow diagram for the present higher-level computer program (the elliptic code). It indicates the sequence of operations, the inter-connections between the various parts of the program, and briefly indicates their function.
PART III

AN INTERMEDIATE-LEVEL CALCULATION METHOD
CHAPTER 4
BASIS OF THE CALCULATION PROCEDURE

4.1 INTRODUCTION:

Unfortunately the "high-level" computer-based flow model which is described in part II above, is unsuitable for providing direct input data into the current generation of building thermal models. This is because the model is very costly in terms of computer time and storage, and is rather user "unfriendly," due to its complexity. In order to overcome this practical limitation, an "intermediate-level" calculation procedure have been developed by Alamdari and Hammond (ref 20), and incorporated into a computer code which the authors have called the ROOM-CHT (Room Convective Heat Transfer) program. This program was envisaged as a "bolt-on" package to building thermal models, and is outlined below.

4.1.1 Requirements of Building Thermal Modellers:

The present generation of building thermal models require surface-averaged convection coefficients for individual room elements: ceilings, floors, walls and windows. Many employ a relatively large time step (~ 60 minutes) and also need these coefficients to be time-averaged over the heating system control cycle. In any case, computing cost would usually prohibit small time step (~ 5 minutes) in design studies even for those models capable of using them (see, for example, Waters (ref 6)). Thus, the normal requirement for multiple averaging suggests that detailed variations in heat transfer distributions across surfaces will have only a minor influence on the final coefficient. This implication of the averaging process lies behind the development of the ROOM-CHT program.

4.2 PHYSICAL BASIS OF THE CALCULATION METHOD:

4.2.1 Profile Analysis of Wall-jets:

Wall-jets are the normal means of air distribution in buildings with forced convective heating systems. The characteristics of these jets are similar to those of classical plane wall-jet in "stagnant" surroundings over much of their development. In order to calculate forced convective heat transfer the present method makes "informed" estimates of the flow and thermal field based on two and three-dimensional turbulent wall-jet characteristics. These jets are assumed to spread out
from their supply apperture and sequentially flow over all room elements. The mean-flow properties for the two-dimensional wall-jet are calculated from the empirical data reviewed by Hammond (ref 33), while that for its three-dimensional counterpart are taken from Rajaratnam (ref 64). Once the mean-flow field has been determined, the corresponding heat transfer distributions across the surfaces are calculated from the "optimum log-law" for wall-jet heat transfer derived by Hammond (ref 34) on the basis of profile analysis.

4.2.1.1 Two-dimensional Wall-jets:

a) Velocity profile:

Coles' (ref 65) velocity profile expressions for plane wall jets in the form used by Hammond (ref 33,66) is adopted here:

$$u^+ = \frac{1}{k} \ln \left( \frac{y^+}{B + C \omega} \right)$$

(4.1)

Where

$$u^+ \equiv \frac{U}{U_T}, \; y^+ \equiv \frac{U_T y}{\nu}, \; b^+_u \equiv \frac{U_T b_u}{\nu}$$

and

$$U_T \equiv (\tau_w/\rho)^{\frac{1}{2}}$$

(4.2)

The velocity half-width, $b_u$, and other wall-jet properties are defined in fig 4.1(a). The values recommended by Brederode and Bradshaw (ref 67) for the log-law constants were adopted by Hammond (ref 34), and have also been used in the present work. These values are: $k=0.41$ and $B=5.2$.

Equation (4.1) is essentially the conventional "log-law" for near wall turbulent shear layers modified by a wake component, $\omega(y^+/(B+b_u))$, and is valid outside the viscous sublayer. Hammond (ref 34) also developed a more complex, two-part expression for the complete velocity profile extending from the wall to the free edge of the jet. However, the variation of velocity within the viscous sublayer is unimportant for the present purposes.
By setting $U^+ = 0$ and $\omega(Y^+/b^+_u) = 1$ at $Y^+ = \delta^+$ (see fig 4.1a) in equation (4.1), the wake function coefficient becomes:

$$C = -\left[ \frac{1}{\kappa} \ln \left( D \ b^+_u \right) + B \right]$$

(4.3)

In the present work the value of the width ratio $D \equiv \delta^+/b^+_u = 2.5$ was employed basis on the profile analysis of Hammond (ref 34).

The velocity profile equation 4.1 may be transformed at its maxima where $Y^+ = Y^+_m$ by noting that,

$$C_f \equiv \frac{\tau_w}{(\frac{1}{2} \rho \ u^2_m)}$$

$$U^+_m \equiv U_m/U_\tau = (2/C_f)^\frac{1}{2}$$

$$Y^+_m \equiv R_m (C_f/2)^\frac{1}{2}$$

(4.4)

This yields a "log-law" for skin friction (ref 34) in the form:

$$\left( \frac{2}{C_f} \right)^\frac{1}{2} = \frac{1}{\kappa} \ln \left[ R_m \left( \frac{C_f}{2} \right)^\frac{1}{2} \right] + B + C \ \omega \left( \frac{Y^+_m}{b^+_u} \right)$$

(4.5)

The value of the wake-component at the velocity peak may be written for convenience in the form of a power-law relation:

$$\omega \left( \frac{Y^+_m}{b^+_u} \right) = 0.205 \ R_m^{-0.1067}$$

(4.6)
A consequence of the solution procedure is that:

\[ b_u^+ = 0.283 R_m \]  \hspace{1cm} (4.7)

b) Temperature Profile:

Similar expression to equation 4.1 may be obtained for the wall-jet temperature profile, outside the molecular diffusive sublayer:

\[ T^+ = \frac{1}{\kappa_\theta} \ln Y^+ + B_\theta^+ + C_\theta \omega_\theta \left( \frac{y^+}{b_\theta^+} \right) \]  \hspace{1cm} (4.8)

where:

\[ T^+ \equiv \frac{T - T_w}{T_f} \]
\[ b_\theta^+ \equiv U_T b_\theta / \nu \]
\[ T \] is the time-averaged value of the absolute temperature,
\[ T_w \] is the wall temperature,
\[ T_f \] is the so-called 'friction temperature' \( \{q_w/(\rho C_p U_f)\} \)
\[ q_w \] is the wall heat flux,
\[ C_p \] is the specific heat of the fluid,

The temperature half-width, \( b_\theta \), and other wall-jet properties are defined in fig 4.1(b). The values recommended by Kader and Yaglom (ref 68) for the log-law constants were adopted by Hammond (ref 34), and have also been used in the present work. These imply a value for turbulent Prandtl number, \( \sigma_t = \kappa / \kappa_\theta \) of 0.85, giving \( \kappa_\theta = 0.48 \). The molecular prandtl number, \( \sigma_\infty \), dependence of the corresponding "additive constant" is given by (ref 68):

\[ B_\theta = 12.5 \sigma_\infty^{\frac{2}{3}} - 5.8 \]  \hspace{1cm} (4.9)
By setting $T^+ = 0$ and $\omega_\theta (Y^+/b^+\theta) = 1$ at $Y^+ = \delta^+\theta$ see fig. 4.1b) in equation (4.8) the thermal wake component coefficient becomes:

$$C_\theta = \left[ \frac{1}{\kappa_\theta} \ln (D_\theta b^+\theta) + B_\theta \right]$$

(4.10)

where the thermal width ratio $D_\theta = \delta^+\theta/b^+\theta$ was given the same value as $D$ by Hammond (ref 34).

The wall-jet temperature profile expression, equation 4.8, may also be transformed at its maxima, where $Y^+ = Y^+m\theta (\Xi U_T Y^+m\theta/\nu)$, by noting that:

$$S_t = \frac{q_w}{\rho C_p U_m (T_m - T_w)}$$

$$T^+_m = \frac{T_m - T_w}{T_T} = \frac{1}{S_t} \left( \frac{C_f}{2} \right)^{1/2}$$

$$Y^+_m \theta = R_m \left( \frac{C_f}{2} \right)^{1/2} \left( \frac{Y_m\theta}{Y_m^+} \right)$$

(4.11)

This leads to Hammond's optimum log-law for heat transfer:

$$\frac{1}{St} \left( \frac{C_f}{2} \right)^{1/2} = \frac{1}{\kappa_\theta} \ln \left[ R_m \left( \frac{C_f}{2} \right)^{1/2} \frac{Y_m\theta}{Y_m^+} \right] + B_\theta + C_\theta \omega_\theta \left( \frac{Y_m\theta}{b^+\theta} \right)$$

(4.12)

Equation (4.12) may be rearranged (ref 34) to yield a complicated relation for the Reynolds analogy factor, $St/0.5 C_f$). This is accomplished by elimination of the terms $R_m (C_f/2)^{1/2}$ in equations (4.5) and (4.12), to give:

$$\frac{St}{0.5 C_f} = \frac{\kappa_\theta / \kappa}{1 + K_\theta \left( \frac{C_f}{2} \right)^{1/2}}$$

(4.13)
The term $K_\theta$ is dependent on the molecular Prandtl number and, according to profile analysis, is given by:

$$
K_\theta = \frac{1}{\kappa} \left[ \ln \left( \frac{Y^{+m\theta}}{Y^{+m}} \right) + B^\theta - B + \kappa_\theta \omega \left( \frac{Y^{+m\theta}}{b^+\theta} \right) \right] - \kappa \omega \left( \frac{Y^{+m}}{b^+u} \right) \] \tag{4.14}
$$

where, $B^\prime = \kappa B$ and $B^\prime_\theta = \kappa_\theta B_\theta$

Recently, Hammond (private communication, 1982) noted that $K_\theta$ was relatively independent of $R_m$, and that it could be more simply represented by:

$$
K_\theta = -11.26 + 14.21 \sigma_\lambda^{0.673} \tag{4.15}
$$

This form of $K_\theta$ has been used in the ROOM-CHT program, although for air ($\sigma_\lambda = 0.71$) $K_\theta$ has a constant value equal to about 0.025. The variation in the profile peak ratio and thermal wake component may be approximated by:

$$
\frac{Y^{+m\theta}}{Y^{+m}} = 1.194 \sigma_\lambda^{-0.455} R_m (0.00209 \sigma_\lambda) \tag{4.16}
$$

$$
\omega \left( \frac{Y^{+m\theta}}{b^+\theta} \right) = 0.169 \sigma_\lambda^{-0.973} R_m (0.0894 \sigma_\lambda^{-0.274}) \tag{4.17}
$$

and for air become:

$$
\frac{Y^{+m\theta}}{Y^{+m}} = 1.395 R_m^{0.0015} \tag{4.18}
$$
\[
\omega_\theta \left( \frac{Y^{+\theta}}{b^{+\theta}} \right) = 0.236 \, R_m^{-0.0982} 
\]  

(4.19)

The analogy factor is insensitive to the choice of the value for the flow specific parameters \( \delta^{+\theta}/b^{+\theta} \) and \( b^{+\theta}/b^{+u} \). In the present work, Hammond’s value of \( b^{+u}/b^{+\theta} = 0.85 \) (ref 34) has been adopted.

In order to convert the optimum log-law in terms of the jet’s initial conditions, it is necessary to make estimates of the mean flow properties of the jet particularly the peak velocity and temperature decay, and the jet spread. These are conventionally written in the form:

\[
\frac{U_m}{U_e} = K_u \left( \frac{x}{S_i} \right)^{-n} 
\]  

(4.20)

\[
\frac{T_m - T_e}{T_e - T_a} = K_t \left( \frac{x}{S_i} \right)^{-n} 
\]  

(4.21)

and

\[
\frac{b_u}{S_i} = K_b \left( \frac{x}{S_i} + \frac{x_0}{S_i} \right) 
\]  

(4.22)

and the local Reynolds number can be expressed as:

\[
R_m = 0.215 \, R_e^{0.89} \left( \frac{x}{S_i} \right)^{0.40} 
\]  

(4.23)

where \( U_e, T_e \) and \( R_e \) are velocity, temperature and Reynolds number at inlet respectively.

The spread constant, \( K_b \), is equal to the far-field spreading rate \( (d b_u/dx) \) and \( x_0/S_i \) is so-called 'geometric' virtual
origin of the jet (ref 69). The variability observed in the experimental data reviewed by Hammond (ref 34, also private communication, 1982) does not seem to justify a choice for the velocity and temperature decay exponent, \( n \), other than 0.5. Adopting these exponents, permits the constants \( K_u, K_t, K_b \) and \( x_0/S_i \) to be assigned mean values from the data in the experimental literature. These constants are given in table 4.1.

<table>
<thead>
<tr>
<th>( K_u )</th>
<th>( K_t )</th>
<th>( K_b )</th>
<th>( x_0/S_i )</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.73</td>
<td>4.09</td>
<td>4.03</td>
<td>8.10</td>
</tr>
</tbody>
</table>

### 4.2.1.2 Three-dimensional Wall-jet:

#### a) Velocity profile

The experimental data reviewed by Rajaratnam (ref 64) suggests that the centreline velocity decay in three-dimensional wall-jet may be expressed in the form:

\[
\frac{U_m}{U_e} = 8.325 \left( \frac{x}{\sqrt{Ag}} \right)^{-1}
\]

(4.24)

where the outlet area, \( Ag \), is the product of its height, \( H_g \), and width, \( W_g \) (see also fig 4.2) and \( U_m \) is the maximum velocity in centreline. According to Rajaratnam the spanwise jet spread is well described by the equation:

\[
\frac{b_z}{W_g} = 0.20 \left( \frac{x}{W_g} \right) ^{-1.25}
\]

(4.25)

where \( b_z \) is the spanwise half-width. Inspection of the available data supports a value for the spanwise width ratio:
\[ \frac{\delta_z}{b_z} = 2.50 \]  

(4.26)

Equation (4.25) breaks down for \( x \leq 15 \) Hg and in this region this spread is better represented by (ref 64):

\[ \frac{b_z}{w_g} = 0.12 \left( \frac{x}{w_g} \right) + 0.20 \]  

(4.27)

Thus, the spanwise jet spread \( \delta_z \), may be calculated as follows:

for the far-field:

\[ \delta_z = 2.5 w_g \left( 0.20 \left( \frac{x}{w_g} \right) - 1.25 \right) \]  

(4.28)

for the near-field:

\[ \delta_z = 2.5 w_g \left[ 0.12 \left( \frac{x}{w_g} \right) + 0.20 \right] \]  

(4.29)

The intersection between these two relations actually occurs at \( x/w_g = 18.125 \).

The spanwise velocity distribution may be written in the form (ref 64):

\[ \frac{U_m}{U_{mo}} = \exp \left[ -0.693 \left( \frac{z}{b_z} \right)^2 \right] \]  

(4.30)

and the potential core length as:

\[ x_c = 8.235 \sqrt{Ag} \]  

(4.31)
b) Temperature profile

In the absence of appropriate experimental data for the temperature field, it is assumed the following relations may be applied (Hammond, private communication, 1982):

\[
\frac{\theta_{mo}}{\theta_e} = 9.125 \left( \frac{x}{\sqrt{Ag}} \right)^{-1}
\]

(4.32)

where \( \theta = T - T_a \), and \( T_a \) is ambient temperature.

The spanwise width ratio may be determined by analogy with Hammond's two-dimensional analysis (ref 34), \( b_z/b_{zo} = 0.85 \), and then the temperature profile may be given by:

\[
\frac{\theta_m}{\theta_{mo}} = \exp \left[ -0.693 \left( \frac{Z}{b_{zo}} \right)^2 \right]
\]

(4.33)

in a similar manner to the corresponding velocity profile.

4.3 COMPUTATIONAL PROCEDURE:

The Reynolds analogy formula (4.12) obtained from profile analysis (Hammond, ref 34) gives the dimensionless wall-jet convection coefficient in the form of Stanton number \( \{St \equiv h_m/(\rho C_p U_m)\} \).

\[
St = St(R_m, \sigma_y)
\]

(4.34)

where, \( h_m \), is heat transfer coefficient \( \{ \equiv q_w/(T_m - T_w) \} \).

However, building thermal modellers conventionally use heat transfer coefficients based on the notional room air temperature:

\[
h_c = q_w/(T_r - T_w)
\]

(4.35)
where, $T_r$, is the room temperature. Alamdari and Hammond (ref 20) have noted that, this coefficient is related to that of the wall-jet by:

$$h_c = h_m \frac{T_m - T_w}{T_r - T_w}$$

(4.36)

The wall-jet convection coefficient may locally fall below the value corresponding to buoyancy-driven convection at the same room temperature difference. In these situations the calculation method adopts the appropriate heat transfer coefficient for buoyancy-driven convection.

The calculation procedure employs improved data correlations (Alamdari and Hammond, ref 70, see Appendix A for bounded paper) to obtain the buoyancy-driven convection coefficient in dimensional terms. Adopting a median "film temperature", $T_f = (T_a + T_w)/2$, applicable to naturally-ventilated buildings (say 27°C) allows the buoyancy-driven convection coefficient to be in the form:

$$h_c = \left\{ \left[ a_c \left( \frac{\Delta T}{L_s} \right) \right]^m + \left[ b_c (\Delta T)^q \right]^m \right\}^{\frac{1}{m}}$$

(4.37)

An examination of experimental data for vertical and horizontal surfaces (see Appendix A) suggests an optimal value of the exponent $m$ of 6. The empirical coefficients, $a_c$ and $b_c$ for both vertical and horizontal surfaces are given in table 4.2. In the present work the characteristic length, $L_s$, has been defined as:

- for vertical surfaces $L_s = H_s$
- for horizontal surfaces $L_s = 4 A_s/P_s$

where

- $H_s$ is the height of the surface,
- $A_s$ is the surface area and
- $P_s$ is the perimeter

In the case of stably-stratified convection near horizontal surfaces the following simple correlating equations is used
\[ h_c = 0.60 \left( \frac{\Delta T}{L_s^2} \right)^{1/5} \]  

(4.38)

Table 4.2; Empirical coefficients in the data correlation for \( h_c \)

<table>
<thead>
<tr>
<th>Flow &amp; surface orientation</th>
<th>( a_c )</th>
<th>( b_c )</th>
<th>( p )</th>
<th>( q )</th>
<th>( m )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buoyancy-driven convection over vertical surface</td>
<td>1.50</td>
<td>1.23</td>
<td>1/4</td>
<td>1/3</td>
<td>6</td>
</tr>
<tr>
<td>Buoyancy-driven flow on horizontal surface</td>
<td>1.40</td>
<td>1.63</td>
<td>1/4</td>
<td>1/3</td>
<td>6</td>
</tr>
</tbody>
</table>

Thus the ROOM-CHT program calculates the local heat transfer distribution over the room internal surfaces in the manner indicated above. The equations to be solved are generally explicit, algebraic ones, except for the wall-jet heat transfer 'log-law' which is implicit and therefore require an iterative solution. In order to avoid the possibility of divergence and to obtain a rapid solution, the Newton-Raphson iteration method (ref 71) has been employed for this equation with convergence being achieved in about three iterations. The computation grid typically employs about 10 uniformly-spaced calculation points per metre length of surface. Once the local heat transfer at mesh nodes was computed, the program performs multiple averaging by successive numerical integrations as required.

The ROOM-CHT program was run with a 45 X 27 uniform solution mesh (its local calculation are automatically grid-independent), and needed a corresponding central processor time of 54 seconds on the Cranfield GEC 4085 mini-computer.
4.4 THE FLOW DIAGRAM FOR THE INTERMEDIATE-LEVEL PROGRAM:

The flow diagram for the intermediate-level (ROOM-CFT) program is shown in figure 4.3. The diagram describes the sequence of the operations and the way different parts of program are connected. It also indicates briefly the functions of various call statements.
PART IV

OBSERVATION AND DEDUCTIONS
CHAPTER 5
COMPUTATIONS AND COMPARISONS

5.1 INTRODUCTION:

This chapter, which presents the major contribution of the present work, is devoted to the comparisons between the results of the higher-level method (the elliptic code) and those of the intermediate-level method (the ROOM-CHT program), which were described in detail in Parts II & III respectively. The validation process consists of comparison with the experimental data obtained in a mechanically-ventilated test room by Alamdari, Chan and Hammond (ref 72). The results of the three-dimensional version of the ROOM-CHT program have been compared with the equivalent two-dimensional version. A parametric study is also reported in which the two-dimensional version of the ROOM-CHT program is used to generate internal surface convection coefficients for some of the most common space heating and air-conditioning geometries used in the UK. 96 different configurations/thermal conditions were examined, in all of which the air supply jet emanated from a linear slot beneath the ceiling. The sensitivity of modern building thermal models to input values for the convective coefficients were studied by Alamdari, Hammond and Melo (ref 81), using a well-tested dynamic model developed by the US National Bureau of Standard. A summary of their findings is presented here.

5.2 WALL-JET STARTING PROFILE COMPARISONS:

In order to overcome the problems associated with using a numerical finite-difference method to compute the jet-induced flow in a large room 'supplied from a small slot inlet (see Section 2.6.1), specified 'starting profiles' were employed at a downstream station (see figure 2.1). In this section, the wall-jet empirical data and Hammond's profile analysis results (ref 33,34), which have been used to generate these starting profiles, are compared with the predicted values obtained from a fine-grid elliptic code for the wall-jet region. Some comparisons are also made with the computations of Ljuboja and Rodi (ref 73,74) using a "boundary layer" or "parabolic" code. The latter code incorporated an algebraic stress turbulence model in which the standard K-ε model, Kolmogorov-Prandtl eddy viscosity constant was replaced by an algebraic expression derived from modelled Reynolds stress transport equations. This more complex model retains a wall damping correction to the Reynolds stress pressure-strain model. In the following comparisons of predicted wall-jet development, two versions of elliptic code were used, one with the standard K-ε turbulence model and the other with an algebraic stress model, similar to that of Ljuboja and Rodi (73,74).
The inlet section of room was modeled (see figure 2.1) with boundary conditions similar to the ones described in Sections 2.6, 2.7 & 3.3, except at the slot inlet, which were prescribed in an analogous manner to that of Pun and Spalding (ref 31) for inlet flow: i.e.

\[
\begin{align*}
U &= U_e \\
V &= 0 \\
H &= H_e \\
K &= 0.00135 U_e^2 \\
\varepsilon &= C_D K_e^{1/2} / (0.03 S_i)
\end{align*}
\]  

(5.1)

The inlet Reynolds number in this particular study was given a value of $7.35 \times 10^3$, with a domain length, $L/S_i=200$, where $L$ is the length of the wall. Predictions were obtained with a non-uniform $31 \times 31$ grid with an expansion ratio of 1.1 in both $x$ and $y$ directions. The converged solution for this case was obtained after about 650 iterations using the standard $K-\varepsilon$ turbulence model, corresponding to a central processor time of about 18000 sec on Cranfield GEC 4085 mini-computer.

5.2.1 Velocity Field:

The streamwise development of the wall-jet velocity field is indicated in figures 5.1 to 5.5. Generally good agreement can be seen between the results of wall-jet profile analysis and empirical data of Hammond (ref 33,34) and turbulence model predictions with both the elliptic code and the parabolic code used by Ljuboja and Rodi. In figure 5.2 the two turbulence models used with the elliptic code are seen to yield velocity profiles that are very close for $y/b_u < 1.30$, but differ slightly nearer the free edge of the jet. By comparison, the profile analysis of Hammond exhibits a "fuller" profile when $y/b_u < 1.0$, and vice versa. The standard $K-\varepsilon$ model version of the elliptic code displays a growth, $d b_u/dx$, that is 15% higher than suggested by empirical data (see figure 5.3), whereas the algebraic stress model predicts a 9% lower rate. The velocity vectors and velocity profiles presented in figure 5.5 are proportional to the local resultant and $x$-direction velocity component respectively, but have not been scaled.
5.2.2 Temperature field:

Figures 5.6-5.8 show the corresponding streamwise development of the temperature field. Although the empirical data for temperature is based on only a very limited number of data sets, reasonable agreement with predictions of the standard K-ε and algebraic stress turbulence models is indicated in figures 5.6-5.8. In a similar manner to the cross-stream profile of velocity (figure 5.2), the profile analysis for the cross-stream temperature profile (figure 5.7) exhibits a fuller profile when y/bο < 1.0 and vice versa. The agreement between the standard K-ε and algebraic stress model predictions for the growth of the wall-jet thermal field, ᵇ/ˢ_i (figure 5.8) is closer than that for ᵇ_u/ˢ_i in figure 5.3. The standard K-ε model gives about 2% lower growth rate than the empirical data suggests, while algebraic stress model yields 7% lower values.

5.2.3 Remarks:

It can clearly be seen in the above comparisons that the difference between the two turbulence models is small in this type of configuration. However, the algebraic stress model required over 100 iterations more than the standard K-ε one. Therefore for the main computations in the whole room, the standard K-ε model of turbulence was adopted.

The close agreement between the wall-jet empirical data reviewed by Hammond, and his profile analysis, with the predictions of both the elliptic and parabolic codes gives confidence in the formers' use. They were therefore employed to specify the starting profiles in the whole room computations.

5.3 COMPARISONS BETWEEN INTERMEDIATE AND HIGH LEVEL METHODS:

5.3.1 Warm-air Heating of Rooms:

5.3.1.1 Choice of simulated operating conditions:

A corner, ground floor domestic living room, having dimensions 4.75m length, 2.45m height and 2.45m width (i.e. L/H=1.94, W/H=1) was modelled (see Fig. 5.9). These dimensions are similar to those used by Sidaway, Hammond and Probert (ref 9) and Sidaway (ref 75). The operating conditions chosen are similar to those used by Alamdari and Hammond (ref 20) to demonstrate the
capabilities of the ROOM-CHT program, and conform with those recommended in the British design manual for gas-fired warm-air heating (ref 76). The two external walls, incorporating single-glazed windows (1.45m X 1.00m in the far-wall and 1.80m X 1.00m in the side-wall), and the floor were given inside surface temperatures estimated on the basis of the best current British practice U-value (ref 20,77). These temperatures are given in table 5.1 as a function of three representative heat loads. A notional room air temperature of 21°C was adopted, while the surface temperature of the internal walls and ceiling were similarly assumed to remain constant at 20°C over the heating season. The commonest terminal device used in warm-air heating practice is the low-side wall register. This was the device studied initially by Alamdari and Hammond (ref 20), and also modelled in the present work. Calculations have been performed for a linear slot spanning the bottom of the wall, with a height of 0.0175m (Si/H=0.007). This is the same slot height as modelled by Alamdari et al (ref 72), Sidaway et al (ref 9) and Sidaway (ref 75). A two-dimensional or plane wall-jet (ref 33,34,64), forms when warm-air is injected into the room. The size adopted for the supply register gives a supply ventilation rate of 8 air-changed per hour (ACH), based on the size of the room and the face velocity (~1.5 m/sec). This is higher than the more realistic domestic ventilation rate of 3.5 ACH adopted by Alamdari and Hammond (ref 20), and is used here only to permit comparison between the two computer codes. The supply conditions for cyclic and modulating control corresponding to the three representative heat loads are given in table 5.2 and 5.3 respectively. The control cycles given for intermittent warm-air heating (see table 5.2) are typical of normal operation (ref 20).

In the case of modulating control, the supply air temperature will modulate very slightly, but, in accordance with Pimbert’s measurements (ref 8), it was maintained constant with time in the calculations at each successive load.

Table 5.1: Demand-dependent Temperatures

<table>
<thead>
<tr>
<th>Demand</th>
<th>Load</th>
<th>Outside Temp. (°C)</th>
<th>Internal Surface Temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Exterior Walls</td>
</tr>
<tr>
<td>High</td>
<td>Full</td>
<td>-1</td>
<td>16</td>
</tr>
<tr>
<td>Intermediate</td>
<td>65%</td>
<td>7</td>
<td>18</td>
</tr>
<tr>
<td>Low</td>
<td>30%</td>
<td>15</td>
<td>20</td>
</tr>
</tbody>
</table>
Table 5.2 ; Supply Conditions - Cyclic Control

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Demand</th>
<th>Supply Air Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Velocity (m/s)</td>
</tr>
<tr>
<td>1</td>
<td>High 20min on/6min off</td>
<td>1.954</td>
</tr>
<tr>
<td></td>
<td>(76.9% on)</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Intermediate 6min on/6min off (50.0% on)</td>
<td>1.954</td>
</tr>
<tr>
<td>3</td>
<td>Low 6min on/20min off (23.1% on)</td>
<td>1.954</td>
</tr>
</tbody>
</table>

Table 5.3 ; Supply Conditions - Modulating Control

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Demand</th>
<th>Supply Air Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Velocity (m/sec)</td>
</tr>
<tr>
<td>1</td>
<td>High</td>
<td>1.503</td>
</tr>
<tr>
<td>2</td>
<td>Intermediate</td>
<td>1.209</td>
</tr>
<tr>
<td>3</td>
<td>Low</td>
<td>0.933</td>
</tr>
</tbody>
</table>

5.3.1.2 Observations:

The predicted flow pattern represented by the resultant velocity vector and the profile for the x-direction component of velocity, and the computed variation in local heat transfer coefficient, \( h_c \), for cyclic and modulating controls corresponding to the three representative heat loads are shown in figures 5.10 to 5.16. The arrows in figure 5.10a represent the resultant velocity vector within the room using the non-buoyant version of elliptic code. The length of the arrows is proportional to the local velocity, but is not scaled. The general shape of the flow pattern (fig 5.10a) and the U-velocity profiles (fig 5.10b) computed using the non-buoyant elliptic code are much the same for all the heating load profiles and control cycles. Only one example of flow pattern is therefore presented here (in Figure 5.10a). It can clearly be seen from figure 5.10a that the jet has spread out from the slot inlet and sequentially flows over the
floor, far-wall and window, ceiling and lastly the near-wall. This is conforms with the assumption made in developing the intermediate-level ROOM-CHT program (ref 20), except that the higher-level elliptic code is able to account for the effect of corners and the flow extract from outlet. The size of the extract slot is not critical in the calculation of the convective heat transfer coefficient (see fig 5.10c), when using the elliptic code, provided that a reasonable number of grid nodes are employed in this region. In the present work the slot outlet’s height was specified to be 0.25 m, which gave 3 grid nodes in slot outlet for 21 x 21 grid with expansion of 1.1 in x and y directions. Figures 5.11a, 5.12a & 5.13a show the flow pattern for the three representative heat load under cyclic control. It is clear from these figures that the reverse flow zone near the window is reduced when the load drops from full to low load. However, this effect is not evident in the case of modulating control (see figures 5.14a, 5.15a, 5.16a), since the supply air velocity in these cases also decreases, and the influence of cold window becomes relatively greater.

The computed heat transfer distribution is plotted on a logarithmic scale in figures (5.11 to 5.16) for clarity. Its variation over the room surfaces is caused by downstream “decay” of wall-jet velocities and temperatures, and by differences in the surface temperature and a notional room air temperature. The higher-level elliptic code is clearly better able to simulate the complicated wall heat transfer distribution resulting from the complex, jet induced room air flow. Nevertheless, the only appreciable difference between the computations of non-buoyant version of the higher-level, elliptic code and intermediate-level, ROOM-CHT program occurs near the corners and the outlet. Although the buoyant version of the elliptic code predicts an appreciably different local variation in $h_c$, particularly on the floor and the ceiling. But in reality, deviations of this magnitude from the values computed by ROOM-CHT program will not have any significant effect on the predictions of dynamic building thermal models. The dependence of the overall, room averaged convection coefficient, $h_{c,r}$ on the heat load for cyclic and modulating controls are indicated in figures 5.17 and 5.18 respectively. In the calculation of this room average convection coefficient, the values given in tables 5.4-5.6 were used. The local values on the side walls were calculated using the improved data correlations for buoyancy-driven convection (ref 70, Appendix A). In cyclic control, the calculated $h_{c,r}$ from ROOM-CHT program is slightly lower than that from the non-buoyant version of the elliptic code, and higher for the buoyant version (see figure 5.17). However, under modulating control the ROOM-CHT value is higher than both versions of elliptic code. The percentage difference between the calculated $h_{c,r}$ from the elliptic code and that from the ROOM-CHT program for both cyclic and modulating controls is shown in tables 5.7 and 5.8.
Table 5.4: Internal Surface Convection Coefficients -- Full Heat Load

\( \frac{W}{m^2 \cdot \kappa} \)

<table>
<thead>
<tr>
<th>FULL HEAT LOAD</th>
<th>SURFACE ELEMENT</th>
<th>Floor</th>
<th>Far-wall &amp; Window</th>
<th>Ceiling</th>
<th>Near-wall &amp; Window</th>
<th>Side-wall</th>
<th>Side-wall</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cyclic Control</td>
<td>ROOM-CHT Program</td>
<td>33.102</td>
<td>5.927</td>
<td>16.023</td>
<td>11.253</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Elliptic Code</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>non-buoyant</td>
<td>33.515</td>
<td>5.592</td>
<td>17.046</td>
<td>17.341</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>buoyant</td>
<td>29.225</td>
<td>3.656</td>
<td>25.570</td>
<td>17.150</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Modulating Control</td>
<td>ROOM-CHT Program</td>
<td>27.359</td>
<td>4.956</td>
<td>13.142</td>
<td>9.225</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Elliptic Code</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>non-buoyant</td>
<td>27.406</td>
<td>4.315</td>
<td>12.888</td>
<td>13.263</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>buoyant</td>
<td>22.170</td>
<td>3.613</td>
<td>25.246</td>
<td>18.139</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SURFACE ELEMENT</td>
<td>FLOOR</td>
<td>Near-wall</td>
<td>Side-wall &amp; Window</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-----------------</td>
<td>-------</td>
<td>-----------</td>
<td>-------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ceiling</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FLOOR</td>
<td>51.398</td>
<td>16.023</td>
<td>11.259</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>COMP-CRT Program</td>
<td>52.331</td>
<td>8.142</td>
<td>18.996</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Elliptic Code</td>
<td>46.356</td>
<td>4.654</td>
<td>20.255</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>buoyant</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>28.982</td>
<td>4.850</td>
<td>6.285</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>COMP-CRT Program</td>
<td>28.826</td>
<td>4.095</td>
<td>8.499</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Elliptic Code</td>
<td>23.536</td>
<td>3.552</td>
<td>17.645</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>buoyant</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 5.5: Internal Surface Convection Coefficients - Intermediate Load

(\text{W/m}^2\text{K})
Table 5.6: Internal Surface Convection Coefficients - Low Load
\( (\text{w/m}^2\cdot\text{K}) \)

<table>
<thead>
<tr>
<th>LOW HEAT LOAD</th>
<th>SURFACE ELEMENT</th>
<th>Floor</th>
<th>Far-wall &amp; Window</th>
<th>Ceiling</th>
<th>Near-wall &amp; Window</th>
<th>Side-wall &amp; Window</th>
<th>Side-wall</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Cyclic Control</td>
<td>ROOM-CHT Program</td>
<td>142.878</td>
<td>18.712</td>
<td>16.023</td>
<td>11.259</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Elliptic Code</td>
<td>146.409</td>
<td>20.219</td>
<td>20.942</td>
<td>19.896</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>non-buoyant</td>
<td>132.673</td>
<td>9.852</td>
<td>24.596</td>
<td>18.272</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>buoyant</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Modulating Control</td>
<td>ROOM-CHT Program</td>
<td>36.331</td>
<td>5.267</td>
<td>4.385</td>
<td>3.142</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Elliptic Code</td>
<td>35.823</td>
<td>4.247</td>
<td>3.993</td>
<td>3.898</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>non-buoyant</td>
<td>28.982</td>
<td>3.350</td>
<td>8.284</td>
<td>7.414</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>buoyant</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: The values are rounded for simplicity.
<table>
<thead>
<tr>
<th>Demand</th>
<th>non-buoyant</th>
<th>buoyant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full</td>
<td>12% higher</td>
<td>6% lower</td>
</tr>
<tr>
<td>Intermediate</td>
<td>11% higher</td>
<td>8% lower</td>
</tr>
<tr>
<td>Low</td>
<td>9% higher</td>
<td>30% lower</td>
</tr>
</tbody>
</table>

Table 5.8: Percentage difference in $\bar{hc,r}$ computed by the elliptic code compared to the ROOM-CHT program - Modulating control

<table>
<thead>
<tr>
<th>Demand</th>
<th>non-buoyant</th>
<th>buoyant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full</td>
<td>18% lower</td>
<td>36% lower</td>
</tr>
<tr>
<td>Intermediate</td>
<td>4% lower</td>
<td>20% lower</td>
</tr>
<tr>
<td>Low</td>
<td>1% lower</td>
<td>22% lower</td>
</tr>
</tbody>
</table>

5.3.2 Air-conditioning of Commercial Offices:

5.3.2.1 Choice of simulated operating conditions:

A room with the same shape and size as that used above for the heating studies, was again employed when examining the convective heat exchange induced by the air conditioning of commercial offices. However, a high-side wall register was adopted for the terminal device, as these are very common in air-conditioning applications. This arrangement is similar to that studied by Sidaway et al (ref 9), Sidaway (ref 75) and Holmes (ref 78). Two different types of structure were chosen for the exterior walls and windows:
Type 1: Cavity wall + Single glazed window
Type 2: Insulant-filled cavity wall + Double glazed window

The U-values corresponding to the above structures are given in table 5.9, (ref 3):

Table 5.9 ; U-values

<table>
<thead>
<tr>
<th>Type of structure</th>
<th>U (W/m² K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single glazed window</td>
<td></td>
</tr>
<tr>
<td>summer</td>
<td>5.90</td>
</tr>
<tr>
<td>winter</td>
<td>6.20</td>
</tr>
<tr>
<td>Double glazed window (13mm)</td>
<td></td>
</tr>
<tr>
<td>summer</td>
<td>3.20</td>
</tr>
<tr>
<td>winter</td>
<td>2.80</td>
</tr>
<tr>
<td>Cavity wall</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.53</td>
</tr>
<tr>
<td>Cavity wall (insulant-filled)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.50</td>
</tr>
<tr>
<td>Floor (ground floor + carpet)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.84</td>
</tr>
</tbody>
</table>

The inside surface temperatures were calculated on the basis of a heat balance, assuming that the outside summer air temperature was 30°C and the inside room air temperature was maintained at 23°C. The surface temperatures corresponding to the structures mentioned earlier are given in table 5.10.

Table 5.10 ; Surface temperature in summer condition (C).

<table>
<thead>
<tr>
<th>Structure</th>
<th>Ceiling &amp; Near-wall</th>
<th>Far-wall</th>
<th>Window</th>
<th>Floor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type 1</td>
<td>23.2</td>
<td>24.3</td>
<td>28.0</td>
<td>23.7</td>
</tr>
<tr>
<td>Type 2</td>
<td>23.2</td>
<td>23.4</td>
<td>25.7</td>
<td>23.7</td>
</tr>
</tbody>
</table>

In air conditioning applications the temperature difference between the supply air at inlet and the return air at outlet is
typically less than 10 C (ref 26). In the present study, the difference between supply air and room air temperatures was assumed to be 12 C in summer season with the corresponding supply air velocity assumed to be 1.75 ms. This velocity was also used as the reference velocity for determining the ratio of maximum to minimum flowrate in cases where "dumping" occurs according to the air-conditioning guide tables (ref 78).

5.3.2.2 Observations:

The predicted flow pattern illustrated by velocity vector plots, U-velocity profiles, and the computed convective heat transfer distribution $hc$, for each structure specified in sub-section 5.3.2.1, are plotted in figures 5.19 to 5.20. A comparison of figures 5.19a and 5.20a, shows that the reverse flow zone near the window is due to the influence of the warm window. It disappears when the single glazed window replaces the double glazed window. Good agreement can be observed for the computed convective heat transfer coefficient predicted by all three programs, particularly for the structure type 2 (see figures 5.19c and 5.20c). The agreement between the two versions of the elliptic codes is very close as shown in figure 5.20c simply because the predicted flow fields (see figure 5.20a) are similar, which in turn is like that assumed for ROOM-CHT program (ref 20).

5.4 EXPERIMENTAL VALIDATION:

In this section, the results of the higher-level and intermediate-level computer programs are compared with the measurements of Alamdari et al (ref 72) for a mechanically-ventilated room.

5.4.1 Operating Conditions:

Alamdari et al (ref 72) have carried out a series of tests in a full-scale environmental test room at Cranfield. The effective internal dimensions of the room were, 4.52m length, 2.52m height and 3.58m width. The terminal device was a high-side wall linear slot register with a height of 0.0175m, and the return air was extracted through a rectangular near-floor outlet. The supply air was provided by a variable speed fan in continuous operation. The measured temperatures are given in table 5.11 for various ventilation rates ranging from 3.3 to 7.65 ACH.
Table 5.11: Measured Surface Temperatures

<table>
<thead>
<tr>
<th>ACH</th>
<th>Ceiling</th>
<th>Far-wall</th>
<th>Floor</th>
<th>Near-wall</th>
<th>Side-walls</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.30</td>
<td>22.5</td>
<td>20.4</td>
<td>19.2</td>
<td>22.4</td>
<td>21.4</td>
</tr>
<tr>
<td>3.99</td>
<td>20.1</td>
<td>18.8</td>
<td>17.0</td>
<td>20.2</td>
<td>19.4</td>
</tr>
<tr>
<td>5.24</td>
<td>23.9</td>
<td>22.5</td>
<td>21.2</td>
<td>24.0</td>
<td>23.1</td>
</tr>
<tr>
<td>6.80</td>
<td>23.1</td>
<td>22.1</td>
<td>21.1</td>
<td>23.6</td>
<td>22.8</td>
</tr>
<tr>
<td>7.65</td>
<td>20.5</td>
<td>19.7</td>
<td>19.3</td>
<td>20.4</td>
<td>19.9</td>
</tr>
</tbody>
</table>

The supply, return and room air temperatures are given in table 5.12.

Table 5.12: Supply and Room Air Temperatures (°C)

<table>
<thead>
<tr>
<th>ACH</th>
<th>Room air</th>
<th>Supply air</th>
<th>Return air</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.30</td>
<td>22.9</td>
<td>26.5</td>
<td>21.3</td>
</tr>
<tr>
<td>3.99</td>
<td>20.4</td>
<td>24.2</td>
<td>19.5</td>
</tr>
<tr>
<td>5.24</td>
<td>23.9</td>
<td>26.9</td>
<td>23.6</td>
</tr>
<tr>
<td>6.80</td>
<td>23.2</td>
<td>25.2</td>
<td>23.1</td>
</tr>
<tr>
<td>7.65</td>
<td>20.5</td>
<td>22.2</td>
<td>20.5</td>
</tr>
</tbody>
</table>

5.4.2 Observations:

Comparisons between the results of the ROOM-CHT program and those of the non-buoyant elliptic code for the mechanically-ventilated test room (ref 72) are shown in figures 5.21 to 5.25.

A typical predicted flow field is presented in figures 5.25b and 5.25c for a mechanical ventilation rate of 7.65 ACH. The dependence of the overall, room-averaged convection coefficients on the mechanical-ventilation rate is shown in figure 5.26, where the calculations of the two computer codes are compared with the measurements of Alamdari et al (ref 72). The large uncertainty band on the experimental data, calculated by the method of Kline and McClintock (ref 80) at 20:1 odds, reflects the problem of
measuring the relatively small temperature differences found in "real" buildings. Calculations of the ROOM-CHT program with and without the buoyancy-driven convection routine are included in figure 5.26. A good agreement between the latter and the elliptic code can be seen in all figures particularly in figures 5.24 and 5.25 for ventilation rates of 6.8 and 7.65 ACH (see tables 5.11 and 5.12).

5.5 COMPARISON BETWEEN TWO- AND THREE-DIMENSIONAL VERSIONS OF THE ROOM-CHT PROGRAM:

The two- and three-dimensional versions of intermediate-level ROOM-CHT program are compared below in order determine whether realistic results could be obtained by assuming a simplified two-dimensional geometry.

5.5.1 Choice of Simulated Operating Conditions:

The geometry and operating conditions for comparison between the 2D and 3D versions of intermediate-level (ROOM-CHT) computer programs are similar to that used when modelling the warm-air heating of rooms (Section 5.3.1), except that, the calculations have been performed for 3.4 air changed per hour and a rectangular grille (200mm X 120mm, 70% free area) at the bottom centre of the interior wall, and also for equivalent linear slot (7.3mm height) spanning the bottom of the wall. These are similar conditions to those reported by Alamdari and Hammond (ref 20).

5.5.2 Observations:

The forced convective surface distribution generated by the three-dimensional wall-jet are shown in figure 5.27. This gives a dramatic illustration of high local coefficients near the supply aperture, which falls off rapidly away from the register. The effect of averaging such local distributions is shown in figures 5.28 and 5.29 for cyclic and modulating controls, corresponding to the three representative heat loads respectively, where the three-dimensional results have been span-averaged, and compared with the equivalent two-dimensional wall-jet. The three-dimensional flow coefficients have a more uniform variation along the floor, and are substantially below the two-dimensional flow values. The disparity in near-aperture heat transfer distribution is, therefore, the major limitation in the using the simpler, two-dimensional geometry.

Only the floor, far-wall/window and ceiling heat transfer distributions are shown in figure 5.27 as the variation over
remaining surfaces is modest, due to the three-dimensional jet having substantially “diffused”. In the surface plot for far-wall and window (fig 5.27b) the convection coefficient over the latter is seen to be lower than the surrounding wall. This is simply due to the way in which the coefficients are factored (see equation 4.36), and the window heat flux itself, \( q_w \), will be higher than that of the surrounding wall.

The time-dependent behaviour induced by cyclic heating control for each surface, and the three representative heat loads, are shown in figures 5.30 and 5.31 for two and three-dimensional ROOM-CHT program respectively. In the two-dimensional version of the program, an overshoot “blip” may be seen on the floor trace (see fig 5.30). This blip displays a very short, but large, convective coefficient associated with the impulsive injection of warm-air. It was calculated on the basis of the experimental data on “starting” wall-jets obtained by Sidaway et al (ref 9). Fortunately, the overshoot has a negligible effect on the time-averaged heat transfer as Alamdari and Hammond (ref 20) first observed, owing to its short duration. The remaining surfaces display a simple rectangular-wave cycle with the dashed line representing the time-averaged coefficients.

The time-averaged heat transfer coefficients are presented in figures 5.32 and 5.33 for cyclic and modulating controls respectively, as a function of the fractional heat load. The rise, at low load, in the convection coefficients for floor and far-wall is another manifestation of their factoring discussed above. The corresponding heat fluxes will in practice be lower, as the temperature differences fall with heat load.

5.5.3 Remarks:

The magnitude of the heat transfer coefficients obtained from three-dimensional version of the ROOM-CHT program are higher than those obtained, for instance, using the CIBS Guide (ref 4) with surface-averaged air velocities calculated via the ROOM-CHT program (see table 5.13). In reality, of course, the designer currently has no means of estimating these velocities (ref 20). The results of the present method are nearer to those that would be expected from the field measurements of Yaneske and Forrest (ref 5), and the building thermal model computational study of Waters (ref 6), although the present results are more detailed.
Table 5.13; Internal Surface Convection Coefficients
(hc, W/m²⁰K)

<table>
<thead>
<tr>
<th>Surface element</th>
<th>Present Method</th>
<th>Cyclic Cont.</th>
<th>Modulating cont.</th>
<th>CIBS Guide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Floor</td>
<td>8.706</td>
<td>4.844</td>
<td>3.825</td>
<td>9.339</td>
</tr>
<tr>
<td>Far-wall</td>
<td>9.031</td>
<td>4.700</td>
<td>3.468</td>
<td>9.758</td>
</tr>
<tr>
<td>Window</td>
<td>1.043</td>
<td>2.572</td>
<td>3.695</td>
<td>1.819</td>
</tr>
<tr>
<td>Ceiling</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Near-wall</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes: Full heat load. *Three-dimensional version

The above version of the ROOM-CHT program yields a notional room-averaged convective heat transfer coefficient at full load of 5.905 and 5.150 W/m²⁰K for cyclic and modulating control respectively. These values are some 40% higher than those originally given by Alamdari and Hammond (ref 20). The main reasons of this increase can be traced to the revised input data, and the adoption of the improved data correlations for buoyancy-driven convection (ref 70, Appendix A). The input data used in the present study differ from that of Alamdari and Hammond (ref 20). In particular, the surface temperature for the internal walls and ceiling were prescribed higher values, and room length was also slightly longer. The present surface internal temperatures were thought to be more realistic, since they ensure that their values never fall below the exterior surfaces temperature no matter what the heat load. In any case, the need to prescribe surface temperatures for example computations such as these is eliminated when the ROOM-CHT program is directly coupled to a building thermal models (see Section 5.7 below), and these temperature are calculated simultaneously.

The CIBS approach (ref 4), obtained in a manner indicated above, gave a coefficient of only 2.182 W/m²⁰K. Yaneske and Forrest (ref 5) have shown that the use of a value as low as the CIBS figure would lead to a substantial under-estimate of the system pre-heat time. The convection coefficients obtained for cyclic control (fig 5.32) are different from those for modulating control (fig 5.33), but the difference is not practically significant. In contrast, the simpler two-dimensional version of the program produced coefficients for floor which are substantially higher (~100%) than its three-dimensional counterpart, for the reasons discussed above. Nevertheless, over all other surfaces, agreement between the two versions of ROOM-CHT program was quite reasonable (ref 20).
5.6 PARAMETRIC STUDIES:

The two-dimensional version of the ROOM-CHT program has been used to generate internal convection coefficients, \( h_c \), for rooms of various dimensions commonly found in domestic and commercial building spaces in the UK. The sizes of the rooms studied here are similar to those used by Holmes (ref 78,79), to demonstrate the application of ceiling mounted air terminal devices for cooling and heating duties. The room lengths and heights were varied from 3m to 5m and 2.5m to 3.25m respectively. In these parametric studies, the size of the window in the far-wall was assumed to be 40% of the room height. A high-side wall, linear slot register, was modelled, from which the supply air was discharged towards the far-wall/window. This is one of the cases examined by Holmes (ref 78,79), in his applications guide. For a given velocity (say 1.75 m s\(^{-1}\)) the height of the slot is made proportional to the size of the room and the number of air changed per hour:

\[
S_i = \frac{HNL}{3600U_e}
\]  

(5.2)

where

- \( N \) is the number of air changed per hour,
- \( H \) is the room height and
- \( L \) is the room length

The supply air temperature in warm-air systems may then be calculated as follows (ref 79):

\[
T_e = \frac{Q}{1000 \rho \dot{V}} + T_r
\]  

(5.3)

where

- \( Q \) is the total design heat load,
- \( \dot{V} \) is the volume flowrate \((=HLW \, N/3600)\)
- \( W \) is the room width

However, for air-conditioning applications the supply temperature is typically 10 \(^\circ\)C below the room air temperature.
The internal surface temperatures were calculated on the basis of a heat balance, with the U-values given in table 5.9, and the room air temperature of 21/23 °C and the outside air temperature of -1/30 °C for the winter/summer seasons respectively. The surface temperatures are given in table 5.14 and 5.15 for winter and summer conditions respectively. Different types of exterior wall and window construction were considered for both summer and winter, as indicated in table 5.16. The internal surface convection coefficients, \( h_c \), are presented in tables B.1 and B.2 in Appendix B.

The computed heat transfer coefficient, \( h_c \), along the near-aperture surface (in this case the ceiling) is substantially higher than those of other surfaces (see tables B.1 & B.2 in Appendix B). These high values of \( h_c \) on the near-aperture surface are due to the way in which the coefficients are factored, for the reasons discussed in Section 5.5. The percentage difference in \( h_c \) between the "standard" case of a single glazed, cavity wall exterior surface (Cases I-III) and Cases IV-VI, VII-IX and X-XII, are indicated in table 5.17, where the effects of having an insulant-filled cavity and double glazing are indicated.

<table>
<thead>
<tr>
<th>Case</th>
<th>Surface element</th>
<th>Ceiling</th>
<th>Far-wall</th>
<th>Window</th>
<th>Floor</th>
</tr>
</thead>
<tbody>
<tr>
<td>I-III</td>
<td></td>
<td>20.5</td>
<td>17.0</td>
<td>4.6</td>
<td>18.8</td>
</tr>
<tr>
<td>IV-VI</td>
<td></td>
<td>20.5</td>
<td>19.7</td>
<td>4.6</td>
<td>18.8</td>
</tr>
<tr>
<td>VII-IX</td>
<td></td>
<td>20.5</td>
<td>17.0</td>
<td>13.6</td>
<td>18.8</td>
</tr>
<tr>
<td>X-XII</td>
<td></td>
<td>20.5</td>
<td>19.7</td>
<td>13.6</td>
<td>18.8</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Case</th>
<th>Surface element</th>
<th>Ceiling</th>
<th>Far-wall</th>
<th>Window</th>
<th>Floor</th>
</tr>
</thead>
<tbody>
<tr>
<td>I-III</td>
<td></td>
<td>23.2</td>
<td>24.3</td>
<td>28.0</td>
<td>23.7</td>
</tr>
<tr>
<td>IV-VI</td>
<td></td>
<td>23.2</td>
<td>23.4</td>
<td>28.0</td>
<td>23.7</td>
</tr>
<tr>
<td>VII-IX</td>
<td></td>
<td>23.2</td>
<td>24.3</td>
<td>25.7</td>
<td>23.7</td>
</tr>
<tr>
<td>X-XII</td>
<td></td>
<td>23.2</td>
<td>23.4</td>
<td>25.7</td>
<td>23.7</td>
</tr>
</tbody>
</table>
Table 5.16: Type of structure

<table>
<thead>
<tr>
<th>Case</th>
<th>Exterior surface</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Wall</td>
<td>Window</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>I-III</td>
<td>Cavity wall</td>
<td>Single glazed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IV-VI</td>
<td>Insulant-filled Cavity</td>
<td>Single glazed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VII-IX</td>
<td>Cavity wall</td>
<td>Double glazed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>X-XII</td>
<td>Insulant-filled Cavity</td>
<td>Double glazed</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 5.17: Percentage difference in $\bar{h}_c$, computed for Cases I-III compared to Cases IV-XII.

<table>
<thead>
<tr>
<th>Condition</th>
<th>% Difference in $\bar{h}_c$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ceiling</td>
</tr>
<tr>
<td>Winter</td>
<td></td>
</tr>
<tr>
<td>Case IV-VI</td>
<td>13% lower</td>
</tr>
<tr>
<td>Case VII-IX</td>
<td>28% lower</td>
</tr>
<tr>
<td>Case X-XII</td>
<td>41% lower</td>
</tr>
<tr>
<td>Summer</td>
<td></td>
</tr>
<tr>
<td>Case IV-VI</td>
<td>Nil</td>
</tr>
<tr>
<td>Case VII-IX</td>
<td>Nil</td>
</tr>
<tr>
<td>Case X-XII</td>
<td>Nil</td>
</tr>
</tbody>
</table>

5.7 SENSITIVITY ANALYSIS ON A BUILDING THERMAL MODEL:

In the context of the work of Alamdari, Hammond and Melo (ref 81) it was felt desirable to quantify the influence on internal and external convection coefficients by using the ROOM-CHT...
and WIND-CHT (another intermediate-level computer code for calculation of external convective heat transfer, ref 81,82) programs respectively on the predicted heating load in a typical domestic dwelling. A hypothetical, detached house was adopted, broadly based on the multi-layered construction of the 3-bedroomed, terraced houses in Livingston, Scotland studied by Clarke and Forrest (ref 83). In addition, a ground floor insulation slab was included (ref 81) to reflect a “heavy” (or thermally “massive”) structure, while the height of each room in the two-storey dwelling were assumed to be 2.8 m. The cavity air gaps and the floor slab were subsequently removed in order to loosely simulate the effect of a “light” structure.

A widely used dynamic model, the NBSLD “response factor” program developed at the US National Bureau of Standard (ref 84), was used to simulate the performance of the above house. The overall response factors contain terms which are sensitive to surface convection coefficients. A large number of discrete hourly intervals are needed to represent such lumped parameters in the case of a building with thermally heavy weight construction, and vice versa (ref 2,81).

In order to simulate the winter heating load Alamdari et al (ref 81), have adopted the meteorological data for Kew, London (51°28’N, 0°19’W) on the 21st December 1965. This constitutes part of the data base for the CIBS “Example Weather Year” (ref 85). An intermittent heating cycle typical of UK practice was assumed (see figure 5.34), with full-load warm-air conditions, similar to those used by Alamdari and Hammond (ref 20). The house has been treated by C. Melo (private communication, 1984) as a five-zone system, with identical thermal conditions in zones adjacent to that being simulated. The ROOM-CHT program was incorporated as an additional subroutine into the NBSLD code, in order to demonstrate the feasibility of direct coupling. The heating load profiles computed by the standard version of the NBSLD program for a city centre location, together with that using the ROOM-CHT program are shown in figure 5.34. The daily energy consumption predicted by the standard program for the heavyweight (lightweight) structure was 382 (432) MJ, and this increased by 7 (11) per cent when the ROOM-CHT code was employed. These computations required corresponding central processor times of 59 and 177 seconds on a Cranfield DEC VAX 11/780 computer, when all five zones were treated sequentially. The heating load profiles shown in this figure display a considerable sensitivity to the internal convection coefficients. The standard version of the NBSLD program employs buoyancy-driven convection coefficients for internal surfaces, and this accounts for the differences in the computed heating load profiles. However, the extend of the impact of the convection model is likely to depend on the conditions prevailing within the particular building being simulated. Nevertheless, the load profile shown in figure 5.34 illustrates the accuracy constraints that are imposed on this building thermal model, and more complex ones, by input internal convection data (ref 81).
CHAPTER 6
CONCLUSIONS AND RECOMMENDATIONS

6.1 CONCLUDING REMARKS:

In an attempt to satisfy the requirements of the new generation building thermal models, a "hierarchy" of interacting and interdependent approaches have been developed for calculating internal surface convective heat transfer coefficients within mechanically-ventilated rooms. These range from "lower-level" approaches, such as improved data correlations for buoyancy-driven convection (Appendix A, ref 70) and wall-jet profile analysis (ref 33,34), to the development of a "high-level" flow model based on finite-difference analogues to the governing "elliptic" equations for the complex, jet-induced room airflow (Part II): the elliptic code. Both the lower and higher-level models have been used to develop and verify an "intermediate-level" computer code (Part III, ref 20,81): the ROOM-CHT program.

In the high-level flow model, "upwind" finite-difference approximation to the governing partial-differential equations are formulated in terms of "primitive" pressure-velocity variables. Closure of these time-averaged, elliptic equations is obtained through an isotropic "eddy" viscosity, or turbulent exchanged coefficient for momentum, calculated via transport equations for both the turbulence kinetic energy and its dissipation rate. The transport of heat is modelled using the effective Prandtl number approach, while "wall functions" are employed to bridge the steep property gradients in near-wall regions. The code of Pun and Spalding (ref 31) was adopted to solve the difference equations for a predetermined size, staggered grid in an iterative "line-by-line" manner using the SIMPLE algorithm of Patankar and Spalding (ref 29).

The ROOM-CHT computer code, employs "informed" estimates of the flow and thermal field based on the known mean-flow properties of wall-jets (ref 33,34,64). The corresponding heat transfer distribution across the room surfaces is calculated using the wall-jet profile analysis, or the improved data correlations for buoyancy-driven convection as appropriate. The program then performs multiple (space and time) averaging by successive numerical integrations as required to meet the needs of building thermal models.

The elliptic code is able to simulate more accurately the complicated wall heat distribution resulting from the complex, jet induced room air-flow. However, this code is rather user unfriendly, since it requires specialist computational thermo-fluids expertise, skill and experience on the part of user. The computational resources for this code are also very high compared with that for the ROOM-CHT program. Alamdari et al
(ref 81) have argued therefore that, the ROOM-CHT program appears to offer the best prospect for meeting the requirements of the new generation of dynamic building thermal models in terms of accuracy, economy and user friendliness. The results of the computations and comparisons of the surface-averaged convection coefficients calculated from ROOM-CHT program, have been shown to be in generally good agreement with those obtained from the elliptic code, and with the limited experimental data. They are also much more realistic than the data currently available to the designer in design guides (see Chapter 5). Nevertheless, ROOM-CHT program for relies its success on the elliptic code which, together with experimental data, provides the basis for its verification. The ROOM-CHT program may therefore be regarded (ref 81) as a means of transferring data in a form appropriate to the needs of building thermal modellers and energy-conscious designers.

6.2 RECOMMENDATIONS FOR FURTHER RESEARCH:

6.2.1 Higher-level Elliptic Code:

The following suggestions for further research should lead to improvements in the accuracy with which the elliptic code is able to predict the flow pattern and surface convective heat transfer within mechanically-ventilated enclosures:

a) The replacement of the upwind finite-difference scheme by Patankar’s power-law scheme (ref 55), which may improve slightly the present results. Although the upwind scheme gives a physically realistic solution, the results of the power-law scheme are much closer to an exact solution, at least for one-dimensional case (see Patankar, ref 55).

b) The rate of convergence might be improved by a change to the SIMPLE algorithm. According to Patankar (ref 55,61), the approximation made in the derivation of P' (see Section 3.2.7), by omission of the terms $\sum a U'$ and $\sum a V'$ from equations 3.40, yields to a rather exaggerated pressure correction. Although this pressure-correction is fairly good at correcting the velocities, it is poor at correcting the pressure. Thus, if the pressure-correction is used only to correct the velocities, and some other means be found to obtain an improved pressure field, this might increase the rate of convergence. Patankar (ref 55,61) has proposed a more efficient algorithm, which he terms "SIMPLER", for this purpose. In this algorithm, the pressure field is obtained from a FDE similar to that for pressure-correction FDE (Eq. 3.45), except that the starred velocities ($U^*$ & $V^*$) in the source term, b, are replaced by the so-called pseudo-velocity ($\tilde{U}$ & $\tilde{V}$), which are defined as:
In this way no approximation is made in the derivation of the pressure equation. Thus, if a correct velocity field is used to calculate the pseudo-velocities, the pressure FDE would at once give the correct pressure.

c) In the case of the buoyant version of the elliptic code, an attempt must be made to increase the rate of convergence, since it was found to be very slow in the present computations. Some improvement might be provided by applying an inertial relaxation method similar to that of Ideriah (ref 52,86). The author attempted to apply Ideriah's relaxation method at an early stage in the development of the buoyancy version of the elliptic code. However, this wasn't very successful at the time, although some variation on this approach might prove more fruitful.

d) Wall-jet profile analysis can be used to specify the starting profiles (see Section 2.6.1) for non-buoyant and/or very weakly buoyant flows with good accuracy. However, this approach is not very suitable for strongly buoyant flows. Improvements in this area are most desirable, since it will not only yield greater accuracy, but might also improve the rate of convergence.

e) The present work was initially based on the belief that a simple, two-dimensional room geometry would probably be adequate for building thermal modelling purposes. However, experience with the ROOM-CHT program, which has been developed with two- and three-dimensional versions, has shown that the development of a three-dimensional version of the elliptic code is desirable. As discussed in Chapter 5, the two-dimensional model produces surface heat transfer coefficients which are substantially too high (~100%) over the surface adjacent to supply aperture (see figures 5.28 & 5.29), although they are quite reasonable over the other surfaces. Unfortunately, the value calculated for the near-supply surfaces has a dominant influence on the overall convective exchange in the room. The Cranfield group under G. P. Hammond are currently engaged in determining the convective heat transfer coefficient over the internal surfaces of three-dimensional, mechanically-ventilated enclosures. Thus the surface coefficients will be obtained by using a high-level, computer-based flow model, together with scale-model experimental measurements, in order to extend and further validate the ROOM-CHT program. The high-level flow model will also facilitate a better understanding of the physics of these complex flows, and permit the calculation of the associated thermal comfort.
6.2.2 Intermediate-level Code:

The main components of the ROOM-CHT program, wall-jet profile analysis (ref 34) and buoyancy-driven convection correlations (Appendix A), have themselves been fully verified by comparison with experiment. It was implicit in the development of the ROOM-CHT program that multi-averaged surface coefficients would be insensitive to such effects. Nevertheless, this needs to be directly verified, and empirical corrections incorporated into the program if necessary. Hammond's group at Cranfield have established an experimental research programme aimed at obtaining the necessary data. The Cranfield group also plan to extend the ROOM-CHT program to permit the calculation of radiative exchange within enclosures. This would enable it to be used to investigate the relative importance of convective and radiative exchange at high ventilation rates. It is hoped that the extended ROOM-CHT program will then be used as the primary "vehicle" for generating design data for a wider range of enclosures and air distribution systems. In this way it would seen feasible to meet the requirements of building thermal modellers for internal surfaces heat transfer coefficients over foreseeable future.
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APPENDIX A

IMPROVED DATA CORRELATION FOR
BUOYANCY-DRIVEN CONVECTION IN ROOMS
Summary Improved data correlations have been derived for buoyancy-driven convective heat transfer from the internal surfaces of naturally-ventilated buildings. They cover the full range of laminar, transitional and turbulent airflows, and are based on the mathematical model of Churchill & Usagi (1972). The new correlating equations are presented in a convenient form for incorporating into modern computer programs which simulate the dynamic thermal performance of buildings. They compare favourably with the available experimental data for isolated surfaces, and are shown to be an improvement on the 'standard' correlations recommended in the CIBS Guide. The factors which affect the accuracy of such data correlations, when used for the energy-conscious design of 'real' buildings, are briefly discussed.

Improved data correlations for buoyancy-driven convection in rooms

F. ALAMDARI, MSc and G. P. HAMMOND, MSc, CEng, MIMechE, MInstR

List of symbols

A, B, C empirical coefficients in correlating Equations (3) and (5)
a, b empirical coefficients in correlating Equation (7)
A, area of heat transfer surface m²
Cp, fluid specific heat at constant pressure J/kgK
g, gravitational acceleration m/s²
Gr, Grashof number (= $\rho^2 g \beta \Delta T L^3 \mu^2$)
h, surface-averaged convective heat transfer coefficient W/m²K
k, fluid thermal conductivity W/mK
L, characteristic length of heat transfer surface m
m, n, p, q exponent in correlating Equations (3), (5) & (7)
Nu, Nusselt number (= hL/k)
Pr, Prandtl number (= Cµ/k)
Pg, perimeter of heat transfer surface m
q, convective heat flux W/m²
Ra, Rayleigh number (= Gr Pr)
T, temperature of the quiescent (ambient) air K
Tf, 'film temperature' K
T, 'wall' (surface) temperature K
$\beta$, coefficient of cubic expansion (= $T_{i}^{-1}$ for gases) K⁻¹
$\Delta T$ (=$T_w - T_q$) K
$\rho$, fluid density kg/m³
$\mu$, fluid dynamic viscosity kg/(m s)

1 Introduction

In the last decade it has become increasingly recognised that in order to develop realistic methods for the energy-conscious design of buildings it is necessary to model the dynamic thermal response of the system. These dynamic models normally require computational solution (as noted in the reviews by Clarke¹ and Day²; the latter summarising the efforts of the UK Science and Engineering Research Council to stimulate research in this area, in contrast to the manual-calculation methods used with the traditional steady-state procedures. Unfortunately, the accuracy of the new generation of building thermal models is presently limited by uncertainties in the input data, particularly for air infiltration and convective heat transfer rates (see, for example, the results of the International Energy Agency (IEA) study reported by Irving³). The aim of the present study was therefore to develop improved methods for calculating buoyancy-driven convective heat transfer within naturally-ventilated buildings.

The convective heat flux from a surface may be written, from 'Newton's Law of Cooling', in the form:

$$q_c = h \Delta T$$

(1)

The surface-averaged convective heat transfer coefficient, $h$, may then be combined with the corresponding radiative coefficient to yield an internal 'surface resistance' for the building elements. In the case of buoyancy-driven (sometimes, but misleadingly, termed 'free' or 'natural') convection, $h$ is itself a function of the temperature difference, $\Delta T$, as well as the length of the surface and the physical properties of the convected fluid. Dimensional analysis may be employed to correlate experimental data reflecting this dependence in terms of dimensionless parameters (defined above in the List of Symbols):

$$Nu = Nu(Gr, Pr)$$

(2)

The Nusselt number relation, for fluids with moderate Prandtl numbers (0.7 < Pr < 70), may be closely represented by a 'power-law' of the form:

$$Nu = C Ra^n$$

(3)

The exponent, $n$, is found to be about 1/4 for low Rayleigh numbers typically in the range $10^4 < Ra < 10^6$, which correspond to laminar flow induced by short surface lengths and/or small temperature differences. Conversely, for $Ra > 10^{10}$ transition to turbulent flow occurs, and $n$ asymptotes to a value of about 1/3 (except for stable-stratification near horizontal surfaces where diffusion on a molecular scale persists up to quite high...
Rayleigh numbers, and \( n \) has a value of about 1/5; see Section 2.3 below). Thus, conventional practice is to calculate \( h \), via data correlations of the form of Equation (3), using an exponent of 1/4 for Rayleigh numbers less than about 10* and 1/3 otherwise. This 'two-part' correlation, implying an abrupt transition, has been adopted in the CIBS Guide. It was also used by the present authors to model off-cycle, buoyancy-driven heat transfer in their 'intermediate-level' calculation procedure for warm-air heated rooms. When used with dynamic thermal models the two-part correlation Equation (3) is usually close to its intersection point, due to the relatively large size of typical building elements: ceilings, floors, walls and windows. This point corresponds to the location of maximum error in the calculated heat transfer coefficient, and the sudden change in the exponent there may also give rise to numerical instability in building thermal models. In Section 2 below, a more elaborate set of correlating equations is developed that cover the full range of laminar, transitional and turbulent flows, and avoid the disadvantages of the simpler, 'two part model'. These improved data correlations are presented in a dimensional form that permit the direct calculation of convection coefficients appropriate to buoyancy-induced air movement at conditions typical of those found in the built environment.

2 Development and validation of the improved correlating equations

The problem of obtaining a correlating equation which will fit both laminar and turbulent buoyancy-driven convection is one of a class of problems involving transfer processes in which solutions are known for asymptotically large and small values of an independent variable, namely:

\[
y - A x^p \text{ as } x - 0
\]

\[
y - B x^q \text{ as } x - \infty
\]

where \( x \) and \( y \) are the independent and dependent variables respectively. Fortunately, Churchill & Usagi have developed a general solution for this class of problems in the form:

\[
y = \left[ (A x^p)^m + (B x^q)^m \right]^{1/m}
\]

(4)

where \( m > 0 \) if \( p < q \), and vice versa. They originally applied this formulation to laminar buoyancy-driven convection with \( x = Pr \), and to laminar, mixed (combined buoyancy-driven and forced) convection. In the present case of laminar/turbulent buoyancy-driven convection a solution may be obtained in the form of Equation (4) as:

\[
\text{Nu} = \left[ (A \text{ Ra}^p)^m + (B \text{ Ra}^q)^m \right]^{1/m}
\]

(5)

An examination of experimental data for vertical and horizontal surfaces (see Sections 2.1 to 2.2 below) suggests an optimal value of the exponent \( m \) of 6. The empirical coefficients in Equation (5) were chosen as mean values from the range employed in the 'standard' data correlations for the asymptotic states published in the literature, and these are summarised in Table 1. This equation, together with the latter coefficients, is valid over an extensive range \((10^4 < \text{Ra} < 10^{12})\) which encompasses all the conditions of practical significance for the built environment. The physical properties of the convected fluid, air, in the case of buildings, may be obtained from standard tables such as those given by Mayhew & Rogers. These properties are primarily dependent on temperature, and their values at the so-called 'film temperature', namely:

\[
T_f = \left( \frac{T_a + T_p}{2} \right)
\]

(6)

are normally employed.

In the context of the built environment the physical properties of air do not vary greatly. It is therefore possible, and obviously convenient, to simplify the 'exact' correlating Equation (5) so that the convection coefficient is recovered in a dimensional form. Adopting a median film temperature applicable to naturally-ventilated buildings (say 27°C) this yields:

\[
h_c = \left[ \left\{ a \left[ \frac{\Delta T}{T_f} \right]^p \right\}^m + \left\{ b \left[ \cdot T_f \right]^q \right\}^m \right]^{1/m}
\]

(7)

where the empirical coefficients, \( a \) and \( b \), for both vertical and horizontal surfaces are again given in Table 1. The temperature variations experienced in buildings imply that the convection coefficients obtained from Equations (5) and (7) are unlikely to differ by more than ± 4 per cent. This is small in comparison with the experimental uncertainty in buoyancy-driven convection data which is typically ± 20 per cent.

The experimental data on buoyancy-driven convection reported in the literature were often obtained using fluids, surface lengths and temperature differences which are unlike those found in buildings. It is therefore desirable to check the empirical coefficients in correlating equations from data more appropriate to this case. Thus, the present data correlations, Equations (5) and (7), were validated by comparison with heat transfer measurements from relatively large area surfaces, mainly in air. Only when data for air were found to be too sparse, were selected measurements for water employed. Details of the comparisons for vertical and horizontal surfaces are discussed in Sections 2.1 to 2.3 below. An idealised representation of the resulting buoyancy-induced flow patterns near isolated surfaces are shown in Fig. 1. These diagrams are used below to distinguish the physical nature of each flow pattern. The effect of the room element and flow interactions present in 'real' buildings are briefly discussed in Section 3.

Table 1. Empirical coefficients in the data correlations for Nu and \( h_c \) (Equations (5) and (7)).

<table>
<thead>
<tr>
<th>Flow and surface orientation</th>
<th>A</th>
<th>B</th>
<th>a</th>
<th>b</th>
<th>p</th>
<th>q</th>
<th>m</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buoyancy-driven convection over vertical surfaces</td>
<td>0.58</td>
<td>0.11</td>
<td>1.50</td>
<td>1.23</td>
<td>1/4</td>
<td>1/3</td>
<td>6</td>
</tr>
<tr>
<td>Buoyancy-driven flow on horizontal surfaces</td>
<td>0.54</td>
<td>0.14</td>
<td>1.40</td>
<td>1.63</td>
<td>1/4</td>
<td>1/3</td>
<td>6</td>
</tr>
</tbody>
</table>

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*J. A. Clarke, private communication, 1981.
2.1 Buoyancy-driven convection near vertical surfaces

The flow pattern induced by a hot, vertical plate results from the heated, lower density fluid adjacent to the surface rising upward (see Fig. 1a). Conversely, a cold, vertical plate, where \( T_w < T_a \), induces a downward motion with a heat flux of similar magnitude, but opposite sign. 'Window downdraught' is an obvious example of the latter motion in the context of buildings. The characteristic length scale, \( L \), associated with these vertical surface flows is the height of the plate. Nusselt number data for air, scaled on this basis, are compared with the present correlating Equation (5) in Fig. 2. It is evident that the scatter of this data is small, and that the improved data correlation exhibits a good fit throughout the laminar, transitional and turbulent regimes. Not surprisingly, therefore, the recovered data for \( h_c \), shown in Fig. 3, is similarly well represented by Equation (7). (The data in this figure implies a slightly greater scatter, due to uncertainties in recovering heat transfer coefficients. Such a process inevitably involves an element of subjective interpretation.) The height of walls in domestic houses are typically about 2.5 m, which is covered by the data displayed in Fig. 3. It is to be expected that heat transfer would be influenced by three-dimensional flow or 'side-wall' effects if the height/width ratio of the walls were relatively large. However, this aspect ratio varies between about 0.5 and 1 in houses, so that the flow and heat transfer are likely to be essentially two-dimensional.

The CIBS Guide does not provide data for surface-averaged, buoyancy-induced heat transfer in the practical case of transition to turbulent flow on vertical surfaces. Its laminar flow data also falls 10 per cent below the
asymptotic Nusselt number given by the present correlating Equation (5). This new expression for convection on vertical surfaces, in the form of Equation (7), therefore provides building thermal modellers with a more accurate and convenient basis for their computations.

When the authors developed the present correlating Equations (5) and (7) they were unaware that later work by Churchill and his co-workers had yielded analogous data correlations for buoyancy-driven motion on vertical surfaces. Churchill & Chu developed their formulation for the two asymptotic conditions \( Ra = 0 \) and \( Ra \to \infty \), with an additional requirement that it should reproduce the analytical results of laminar boundary layer theory in the range \( 10^5 < Ra < 10^7 \). This led to the rather complicated expression:

\[
\text{Nu} = 0.825 + \frac{0.387 \cdot Ra^{1.6}}{1 + (0.437/Pr)^{0.618^{0.27}}} \tag{8}
\]

Very low Rayleigh numbers \( (Ra = 0) \) are only encountered in practice for rather exotic low Prandtl number fluids, such as liquid metals, which are obviously inappropriate to the built environment. Nevertheless, ESDU has simplified Equation (8) for air flow and, over the limited range of temperatures applicable to naturally-ventilated buildings, it becomes:

\[
h = 0.134 \cdot L^{-1} + 1.11 \cdot \sqrt{T}^{1/6} \tag{9}
\]

This expression, together with its counterpart, Equation (8), are found to yield heat transfer coefficients which are about 5 per cent higher than the present correlating Equations, (5) and (7), for high temperature differences and long surfaces (high Rayleigh numbers, \( 10^9 < Ra < 10^{12} \)). Conversely, for relatively low temperature differences and short surface lengths \( (10^5 < Ra < 10^7) \), the Churchill & Chu/ESDU equations imply coefficients that are about 11 per cent lower than those given by the present correlations. Interestingly, Churchill & Chu themselves argued that heat transfer coefficients similar to those yielded by the present correlating Equation (5) are 'more accurate', in the range \( 10^5 < Ra < 10^7 \), than those of Equation (8). Nevertheless, in view of the inherent uncertainty in the experimental data on buoyancy-driven convection noted above, the Churchill & Chu/ESDU correlating equations may be regarded as alternatives to the present ones for vertical surface flows. The present data correlations, however, have the additional merit of being rather simpler in form, and have been extended to horizontal surfaces in the section that follows.

2.2 Buoyancy-driven convection near horizontal surfaces

The motion induced by a heated horizontal surface is illustrated in Fig. 1b. The relatively hot, lighter fluid on the upper surface has a tendency to be convected upwards in the form of 'plumes' or 'thermals', being replaced by colder, more dense fluid from above. This configuration is therefore gravitationally unstable and, for temperature differences above a minimum or 'critical' value, the buoyancy forces will drive the convective motion. The flow pattern shown for the upper surface in Fig. 1b is an idealisation based on the work of the Soviet scientist M. Mikheyev as given by Al-Arabi & El-Riedy.

*The authors only discovered a reference to the paper of Churchill & Chu when they consulted the Engineering Sciences Data Unit in search of additional data to validate the horizontal surface cases.

The motion induced by a heated horizontal surface is illustrated in Fig. 1b. The relatively hot, lighter fluid on the upper surface has a tendency to be convected upwards in the form of 'plumes' or 'thermals', being replaced by colder, more dense fluid from above. This configuration is therefore gravitationally unstable and, for temperature differences above a minimum or 'critical' value, the buoyancy forces will drive the convective motion. The flow pattern shown for the upper surface in Fig. 1b is an idealisation based on the work of the Soviet scientist M. Mikheyev as given by Al-Arabi & El-Riedy.

'Orderly', laminar flow is observed at low Rayleigh numbers \( (Ra < 10^6) \), but the convective motion undergoes a transition to turbulence at high values \( (Ra > 10^9) \). A cooled horizontal surface, where \( T_\infty < T_\wedge \), will give rise to a similar flow pattern, although this develops on the underside of the element. Both arrangements involve a downward heat flow. There is a considerable divergence of opinion in the literature regarding the most appropriate length scale to adopt for these horizontal surface flows. In the present study the characteristic length has been defined as:

\[
L = 4A/P_s \tag{10}
\]

where \( A \) is the surface area and \( P_s \) its perimeter. This is analogous to the 'hydraulic diameter' used in fluid mechanics, and for a square surface the length scale simply becomes the length of one side. Equation (10) has the conceptual merit of yielding appropriate length scales for irregular, non-rectangular surfaces, although elements of this type are not usually found in buildings. Experimental Nusselt number data obtained in air and corrected for the present length scale are shown in Fig. 4, where they are compared with the new correlating Equation (5). The data of Fujii & Imura for water (considered by Yousef et al to be amongst the most accurate) has also been included in Fig. 4 in order to augment the rather sparse air data at high Rayleigh numbers. Data for buoyancy-driven convection on horizontal surfaces is seen to display a greater scatter than that for vertical surfaces. This is in part due to the influence of 'edge' effects which tend to increase the heat transfer rate on relatively small surfaces.

Nevertheless, the present correlating Equation (5) is seen to be representative of the bulk of this data throughout the laminar, transitional and turbulent regimes. A similar
Conclusion may be drawn in regard to the recovered data for \( h \), in air plotted on Fig. 5, where they are compared with correlating Equation (7). Yousef et al.\(^1\) postulated that the very high heat transfer coefficients obtained with their smallest test plate (\( L = 0.1 \) m) might have been caused by the presence of a single buoyant plume. This explanation for the anomalous data is open to doubt, but this geometry is, in any case, outside the range of interest to the built environment. The data of Saunders et al.\(^1\) which fall considerably below the present correlation, is generally regarded (see, for example, Yousef et al.\(^1\)) as being among the lowest of the available data sets. It is clear from Fig. 5 that further accurate data for \( L = 3 \) m would have been useful for validating the present correlating Equation (7) at conditions directly applicable to buildings.

Both the CIBS Guide\(^4\) and the appropriate ESDU Data Item\(^3\) employ two-part models to correlate data for \( h \), in air near horizontal surfaces. The former adopts a characteristic length scale equal to the mean value of the two sides of a rectangular surface. This choice is identical to that given by Equation (10) for square surfaces, and differs only slightly for rectangular elements. The length scale adopted by ESDU is equal to \( 0.25 L \) as used in the present work. In the laminar regime, the CIBS correlation is almost identical to that given by Equation (7), whereas the ESDU relation yields coefficients which are almost 30 per cent higher. The CIBS and ESDU correlations for the turbulent regime, in contrast, imply coefficients which are not more than 5 per cent above those given by correlating Equation (7).

2.3 Stably-stratified convection near horizontal surfaces

The fluid beneath a heated horizontal surface, such as that illustrated in Fig. 1b, must always be at a lower temperature than that of the surface. This fluid is therefore gravitationally stable, and the layer over which the vertical temperature gradient persists is stably-stratified. A cooled horizontal surface would also give rise to the formation of such a stable layer, but on top of the element. Either configuration may be characterised as having an upward heat flow. Instabilities that would otherwise lead to the onset of turbulence at high Rayleigh numbers are damped under the influence of downward, negative buoyancy forces which restore equilibrium. The analytical results of Singh et al.\(^1\) based on the solution of the integral boundary layer equations, indicate that the Nusselt number for such laminar stably-stratified layers varies according to a \( 1/5 \) power of the Rayleigh number. This differs from the \( 1/4 \) power-law dependence of buoyancy-driven laminar flow on both vertical and horizontal surfaces. Nevertheless, recent experimental data\(^4,14,15\) has confirmed that the \( 1/5 \) power relation holds up to at least \( Ra = 10^{10} \), and it is adopted in the present work. Thus, for stable stratification alone, the standard correlation Equation (3) is used, together with the \( 1/5 \) exponent and the empirical coefficient, \( C \), having a value of 0.58. Simplifying this expression for the thermal conditions applicable to naturally-ventilated buildings yields a dimensional formula for the convective heat transfer coefficient:

\[
h_c = 0.60 \left( \frac{\Delta T}{L^2} \right)^{1/5}
\]

The choice of the characteristic length scale governing heat transfer across stably-stratified layers has again been the subject of divergent opinions in the literature. Equation (10) has been employed in the present work for the reasons outlined in Section 2.2 above. The experimental Nusselt number data for air\(^4,14,15,19\) which is displayed in Fig. 6 has been scaled using this characteristic length. The early measurements by
Saunders et al., on which the 'standard' data correlations are based, have been criticised by subsequent investigators and are regarded as too high. They used a test plate which simultaneously heated the fluid above and below the surface, so that the flow over the upper surface influenced that near the edges of the lower one. Restrepo & Glicksman have also drawn attention to the influence of the edge thermal boundary conditions (whether they are heated, cooled or insulated) which can alter the heat transfer coefficient by more than \( \pm 20 \) per cent. In order to give fuller coverage of high Rayleigh number data, the measurements of Fujii & Imura for water have been included in Fig. 6. It can be seen that the present correlating Equation (3) is representative of the more recent measurements, and that the early data of Griffiths & Davis and Saunders are appreciably higher. This is the reason that correlating Equation (11) does not appear to fit the recovered data for \( h \), in Fig. 7 very closely. In fact the more recent, but rather sparse, data is fairly evenly scattered about the present data correlation.

A data correlation based on the measurements of Saunders et al. employing a \( \frac{1}{4} \) power-law dependence has been adopted for both the CIBS Guide and the corresponding ESDU Data Item. These ‘standard’ correlations, when corrected for the length scale given by Equation (10), are almost identical, but yield heat transfer coefficients which are up to 40 per cent higher than those given by correlating Equations (3) and (11) under conditions typical of the built environment. Indeed the older correlations are over 120 per cent in error at the highest Rayleigh number of their declared range of application. The use of the \( \frac{1}{4} \) power-law is at variance with theoretical results and with recent experimental data, which all suggest a 1/5 power-law dependence for heat transfer in stably-stratified layers. The new correlating Equations (3) and (11) therefore constitute a significant improvement on these older data correlations.

3 Concluding remarks

Improved data correlations for buoyancy-driven convection from the internal surfaces of naturally-ventilated buildings have been derived on the basis of the model of Churchil & Usagi. These correlating equations provide a smooth fit to data across the full range of laminar, transitional and turbulent airflows, in contrast to the 'two-part' model employed in 'standard' correlations. The new correlating equations thereby avoid the possibility of inducing numerical instability when used in conjunction with the new generation of dynamic building thermal models. The new data correlations are presented in a simplified and more convenient form, Equation (7), from which the convective heat transfer coefficient may be obtained directly in dimensional terms, rather than via the 'exact' Nusselt number relation. They display generally good agreement with the rather limited experimental data available for conditions typical of the built environment. However, it has been shown that the standard, two-part correlation equations used by the CIBS and the ESDU are, in some cases, significantly in error when compared with recent measurements.

The data on which the improved correlating equations are based was obtained from experiments using isolated surfaces. It is therefore appropriate to consider the extent to which this idealisation is valid in the conditions prevailing in 'real' buildings. The ESDU has identified a number of factors which alter the apparent heat transfer rates implied by correlating equations, and those relevant to buildings include non-uniform surface temperatures, room temperature stratification, room element and flow interactions, and draughts. Surface temperatures in naturally-ventilated buildings are reasonably uniform, and vertical room temperature gradients of 1°C per metre (dictated by thermal comfort requirements) would also have a negligible effect on surface heat transfer. However, the effect of room element interactions and draughts are more problematic. The data of Griffiths & Davis demonstrates that the interaction between a floor and an adjacent wall boundary layer is negligible, although that between such a layer and a ceiling (for which there appears to be no direct data available) is likely to be significant. The influence of draughts is similarly uncertain. In addition, heat transfer within rooms will be affected by the presence of fixtures and fittings, as well as surface texture or relative roughness. Fortunately, surface-averaged heat transfer coefficients for the large surface areas associated with building elements are generally insensitive to local variations such as these. Nevertheless, building thermal modellers should allow for variations of at least \( \pm 20 \) per cent in the coefficients recommended for correlating Equations (7) or (11) to account for these disturbances. Despite the obvious limitations in using data for idealised buoyancy-driven convective processes, correlating equations such as those developed here remain arguably the most appropriate calculation method in the absence of field measurements.

Finally, it must be emphasised that the present heat transfer data correlations only apply to buildings that are naturally-ventilated. In order to determine the thermal...
performance of buildings using forced convective heating and/or mechanical ventilation, the convection coefficient must be calculated via a more elaborate procedure, such as that recently developed by the present authors. Yaneske & Forrest concluded for example, from a field study of over thirty rooms of various shapes and sizes heated by a fan 'convectA)r', that surface coefficients were much higher than those for buoyancy-driven convection. They demonstrated that the use of the latter when selecting heater capacity for a mechanically-ventilated dwelling could result in substantial increases in preheat times. Waters has likewise shown that the accuracy of his implicit finite-difference building thermal model was strongly dependent on the correct choice of internal heat transfer coefficient when simulating mechanically-ventilated structures.

Acknowledgements

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The authors' names appear alphabetically.

References

12 ESDU Data Item No. 77031, Heat transfer by free convection and radiation—simply shaped bodies in air and other fluids (ESDU, London, 1979).
APPENDIX B

INTERNAL SURFACE CONVECTION COEFFICIENT

(Results of Parametric Studies)
Table B.1: Internal Surface-averaged Convection Coefficients - Winter Season

<table>
<thead>
<tr>
<th>Winter Case No. I</th>
<th>Room length 3.0 (m)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Single glazed window</td>
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<td></td>
<td>Cavity wall</td>
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</table>

<table>
<thead>
<tr>
<th>Room height (m)</th>
<th>Internal Surface Convection Coefficient (W/m² K)</th>
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<tbody>
<tr>
<td></td>
<td>Ceiling</td>
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<tr>
<td>2.50</td>
<td>87.539</td>
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<td>3.00</td>
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<td></td>
<td>Cavity wall</td>
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<table>
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<th>Room height (m)</th>
<th>Internal Surface Convection Coefficient (W/m² K)</th>
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<td>Cavity wall</td>
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Table B.1: -- continue

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<td>Room height (m)</td>
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</tr>
<tr>
<td>Ceiling</td>
<td>Far-wall</td>
<td>Floor</td>
<td>Near-wall</td>
<td>Window</td>
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<tr>
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<td>Far-wall</td>
<td>Floor</td>
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Winter Case No. VII
- Room length 3.0 (m)
- Double glazed window
- Cavity wall

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Winter Case No. VIII
- Room length 4.0 (m)
- Double glazed window
- Cavity wall

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Winter Case No. IX
- Room length 5.0 (m)
- Double glazed window
- Cavity wall
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Table B.2: Internal Surface-averaged Convection Coefficients - Summer Season

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### Table B.2: -- continue

#### Summer Case No. VII
- Room length 3.0 (m)
- Double glazed window
- Cavity wall

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#### Summer Case No. VIII
- Room length 4.0 (m)
- Double glazed window
- Cavity wall

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#### Summer Case No. IX
- Room length 5.0 (m)
- Double glazed window
- Cavity wall

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FIGURES
Fig. 1.1: World Population & Rate of Energy Consumption: Projections (O’Callaghan, ref 87).
Fig. 2.1: Two-dimensional Room & Starting Wall-jet Profile.
Fig. 3.1: The illustration of the two-dimensional 21x21 grid with expansion of 1.3.
Fig. 3.2: Two-dimensional Control Volume for General Variable $\bar{\phi}$.

Fig. 3.3: Location of the Dependent Variables;

- $\rightarrow$ = U-velocity
- $\uparrow$ = V-velocity
- $\circ$ = Other variables.
Fig. 3.4: Control Volume and Staggered Locations for U.

Fig. 3.5: Control Volume and Staggered Locations for V.

Fig. 3.6: One-dimensional Grid and Control Volume.
Fig. 3.7: Variation of the Coefficient $a_E$ with Peclet number (Patankar, ref 55).
Fig. 3.8: Boundary Control Volume of Variables.

Fig. 3.9: Boundary Control Volume for $\bar{\phi}$ and $U$. 
Fig. 3.10: Representation of Line-by-line Iterative Method.
Fig. 3.11: The Initialized U-velocity Prior to the First Iteration of Initial Run.
Fig. 3.12: Computed Local Heat Transfer Coefficients along the Floor (Grid-dependence test).
Fig. 3.13: Predicted U-velocity Profile (Grid-dependence test).
Fig. 3.14a: Computed Local Heat Transfer Coefficients along the Floor, for 21X21 Grid (Grid expansion test).
Fig. 3.14b: Computed Local Heat Transfer Coefficients with 21X21 Grid and Expansion 1.1.
Fig. 3.14c: Computed Local Heat Transfer Coefficients with 21x21 Grid and Expansion 1.3.
Start

Input data

Preliminaries

Grid specification, reference values, etc...

Initialise dependent variable fields

Start field iteration sweep

Select new traverse line

If Iteration is > X

Set boundary conditions not requiring updating during iterations

Solve for dependent variable Eq 3.17 & 3.38

Update the turbulent viscosity

Solve for pressure correction equation Eq 3.45

Adjust U, V & P for each cell on TDMA line Eq 3.44 & 3.39

Subtract reference pressure from pressure field & set boundary values for print out

If Iteration is \( \leq 1 \)

Calculate FDE coefficient and source terms Eq 3.18 & 3.46

Modify the source terms to apply boundary conditions

Solve FDE by TDMA Eq 3.48 & 3.51

yes

Decide whether to repeat no the line traverse

Decide whether all the yes lines have been visited

Apply block-correction procedure Eq 3.54 & 3.55

no

Decide whether solution has converged

yes

Print out

Stop

Fig. 3.15: Flow-diagram for the High-level Code.
\[ U^+ = \frac{dU^+}{dY^+} = \frac{d^2U^+}{dY^{+2}} = 0 \]

\[ U = 0, \quad \frac{dU^+}{dY^+} = 1 \]

(a)

\[ T^+ = \frac{dT^+}{dY^+} = \frac{d^2T^+}{dY^{+2}} = 0 \]

\[ T = 0.5 T_m, \quad \frac{dT^+}{dY^+} = 0 \]

\[ T = T_m, \quad \frac{dT^+}{dY^+} = 0 \]

(b)

Fig. 4.1: Wall-jet Notation and Boundary Conditions (Hammond, ref 34).
Fig. 4.2: Schematic Representation of Three-dimensional Wall-jet, (Rajaratnam, ref 64).
Start

Input data

Grid specification

Evaluate the corresponding length scale and area of elements

Calculate the Reynolds number at inlet

Call BUOY

Calculate buoyancy-driven convective heat transfer coefficients, using improved data correlations

3D

Decide for 2/3D problem

Calculate averaged convective heat transfer on each element

Call FORCE2

Calculate the local forced convective heat transfer coefficients

2D

Decide for 2/3D problem

Call FORCE3

Evaluate the time-averaged force convective heat transfer coefficients

M

Decide whether cyclic or modulating control

C

Call TIMEAN

Print out

Stop

Fig. 4.3: Flow-diagram for the Intermediate-level (the ROOM-CHT) Program.
Empirical data (Hammond, ref 33)

Parabolic code:
- Algebraic stress model (Ljuboja & Rodi, ref 73)

Elliptic code (present computations):
- Standard K-\(\varepsilon\) model
- Algebraic stress model

Fig. 5.1: Streamwise Development of \(U_m/U_e\).

Profile analysis (Hammond, ref 34)

Elliptic code (present computations):
- Standard K-ε model
- Algebraic stress model

Fig. 5.2: Cross-stream Distribution of U/Um.
Empirical data (Hammond, ref 33, 34)

Parabolic code:
\( \triangle \) Algebraic stress model
(Ljubojca & Rodi, ref 73)

Elliptic code (present computations):
\( \square \) Standard \( K-\varepsilon \) model
\( \circ \) Algebraic stress model

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Fig. 5.3: Streamwise Development of \( b_u/S_i \).
Empirical data (Hammond, ref. 33, 34)

Elliptic code (present computations):
- Standard \( K-\varepsilon \) model
- Algebraic stress model

Fig. 5.4: Streamwise Momentum Distribution.
Fig. 5.5: Wall-jet Predicted Velocity Field.
Empirical data

Elliptic code (present computations):
- Standard K-\( \varepsilon \) model
- Algebraic stress model

Fig. 5.6: Streamwise Development of \((T_m - T_a)/(T_e - T_a)\).
Profile analysis (Hammond, ref 34)

Elliptic code (present computations):
- Standard K-ε model
- Algebraic stress model

Fig. 5.7: Cross-stream Distribution of \((T-T_a)/(T_m-T_a)\).
Empirical data (Hammond, ref 33,34)

Elliptic code (present computations):
- Standard K-\(\varepsilon\) model
- Algebraic stress model

Fig. 5.8: Streamwise Development of \(b_\theta/S_i\).
Fig. 5.3: Schematic Diagram of Room with "Low Side Wall Register."
FIG. 5.10b:
Predicted U-velocity for Case No. 1 of Table 5.2.

Non-buoyant version of the Elliptic Code.
Fig. 5.10c: Computed Local Heat Transfer Coefficients—Test for Slot Outlet’s Height.
Fig. 5.11a: Predicted flow pattern for Case No. 1 of Table 5.2.
Fig. 5.11c: Computed Local Heat Transfer Coefficients for Case No. 1 of Table 5.2.
Fig. 5.12b: Predicted U-velocity profile for Case No. 2 of Table 5.2.
Fig. 5.12c: Computed Local Heat Transfer Coefficients for Case No. 2 of Table 5.2.
Fig. 5.13b: Predicted U-velocity Profile for Case No. 3 of Table 5.2.
Fig. 5.13c: Computed Local Heat Transfer Coefficients for Case No. 3 of Table 5.2.
Fig. 5.14a: Predicted Flow Pattern for Case No. 1 of Table 5.3.
Fig. 5.14c: Computed Heat Transfer Coefficients for Case No. 1 of Table 5.3.
Fig. 5.15a: Predicted Flow Pattern for Case No. 2 of Table 5.3.
Fig. 5.15b: Predicted U-velocity Profile for Case No. 2 of Table 5.3.
Fig. 5.15c: Computed Heat Transfer Coefficients for Case No. 2 of Table 5.3.
Fig. 5.16a: Predicted Flow Pattern for Case No. 3 of Table 5.3.
Fig. 5.16b; Predicted U-velocity Profile for Case No. 3 of Table 5.3.
Fig. 5.16c: Computed Heat Transfer Coefficients for Case No. 3 of Table 5.3.
Fig. 5.17: The Dependence of the Overall, Room Averaged Convection Coefficient on the Heat Load - Cyclic Control.
Fig. 5.18: The Dependence of the Overall, Room Averaged Convection Coefficient on the Heat Load - Modulating Control.
Predicted Flow Pattern for Structure Type 1 of Table 5.10.
Fig. 5.19b: Predicted U-velocity Profile for Structure Type 1 of Table 5.10.
Fig. 51.9c: Computed Heat Transfer Coefficients for Structure Type 1 of Table 5.10.
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Fig. 5.21: Computed Heat Transfer Coefficients for Conditions of 3.3 ACH of Tables 5.11 and 5.12.
Fig. 5.22: Computed Heat Transfer Coefficients for Conditions of 3.99 ACH of Tables 5.11 and 5.12.
Fig. 5.23: Computed Heat Transfer Coefficients for Conditions of 5.24 ACH of Tables 5.11 and 5.12.
Fig. 5.24: Computed Heat Transfer Coefficients for Conditions of 6.8 ACH of Tables 5.11 and 5.12.
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Fig. 5.25b: Predicted Flow Pattern for Conditions of 7.65 mCHH of Tables 5.11 and 5.12.
Fig. 5.25c: Predicted U-velocity Profile for Conditions of 7.65 ACH of Tables 5.11 and 5.12.
Fig. 5.26: Mechanical-ventilation Rate Dependence of the Room-averaged Convection Coefficient.
Fig. 5.27a: Three-dimensional 'Wall-jet' Heat Transfer Distribution along the Floor (hc, W/m K).
Fig. 5.27b: Three-dimensional "Wall-jet" Heat Transfer Distribution along the Far-wall and Window (hc, W/m K).
Fig. 5.27c: Three-dimensional "Wall-jet" Heat Transfer Distribution along the Ceiling (hc, W/m K).
Two-dimensional wall-jet

Three-dimensional wall-jet

Fig. 5.28a: Comparison of Floor Heat Transfer Distribution for Case No. 1 of Table 5.2.
Fig. 5.28b: Comparison of Floor Heat Transfer Distribution for Case No. 2 of Table 5.2.

- Two-dimensional wall-jet
- Three-dimensional wall-jet
Fig. 5.28c: Comparison of Floor Heat Transfer Distribution for Case No. 3 of Table 5.2.
Fig. 5.29a: Comparison of Floor Heat Transfer Distribution for Case No. 1 of Table 5.3.
Two-dimensional wall-jet

Three-dimensional wall-jet

Fig. 5.29b: Comparison of Floor Heat Transfer Distribution for Case No. 2 of Table 5.3.
Fig. 5.29c: Comparison of Floor Heat Transfer Distribution for Case No. 3 of Table 5.3.
Fig. 5.30b: Time-dependent heat transfer for Case No. 2 of Table 5.2 (two-dimensional version).
Fig. 5.30c: Time-dependent Heat Transfer for Case No. 3 of Table 5.2 (Two-dimensional version).
Fig. 5.31a: Time-dependent heat transfer for Case No. 1 of Table 5.2 (three-dimensional version).
Fig. 5.3lb: Time-dependent Heat Transfer for Case No. 2 of Table 5.2 (Three-dimensional version)
Fig. 5.3lc: Time-dependent Heat Transfer for Case No. 3 of Table 5.2 (Three-dimensional version).
Fig. 5.32: Time-averaged Room Heat Transfer Under Cyclic Control (see table 5.2).
Fig. 5.33: Time-averaged Room Heat Transfer Under Modulating Control (see Table 5.3).
Fig. 5.34a  Predicted winter heating load profile for a detached house in the UK.
Fig. 5.34b  Predicted winter heating load profile for a detached house in the UK.