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ANALYSIS OF COMBUSTION INSTABILITY IN LIQUID FUEL ROCKET MOTORS

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FOREWORD

This report summarizes a portion of the work done for NASA Grant NGR 43-003-015. It is the Ph.D. research of the first author, Kin-Wing Wong. John Peddieson was the advisor; M. Ventrice was the principal investigator of the grant.

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Chapter 1

INTRODUCTION

The steady operation of a liquid-propellant rocket engine is often disturbed by the occurrence of large pressure oscillations in the combustion chamber. These oscillations, which can lead to damage to or failure of the motor, are caused by amplification of initially small acoustic disturbances due to the energy released by unsteady burning of the propellant. This is usually referred to as combustion instability. It is generally accepted that unsteady burning can be correlated with the gas pressure (pressure sensitivity) and the magnitude of velocity of the fuel drops relative to the gas (velocity sensitivity). This dissertation is concerned with mathematical modeling of velocity-sensitive combustion instability in liquid-propellant rocket motors.

A survey of literature dealing with mathematical modeling of pressure-sensitive combustion instability can be found in the dissertation of Powell (1). One of the first papers to examine nonlinear effects is that of Maslen and Moore (2) who considered only the fluid mechanical effects in a circular cylinder. The paper by Priem and Guentert (3) is one of the first to include the effect of velocity sensitivity. They discussed combustion instability

in a thin annulus.

Since direct numerical solution of the governing equations is very time consuming and difficult to extract information from, many investigators such as Powell and Zinn (4), Lores and Zinn (5), and Peddiesion, Ventrice, and Purdy (6) have employed methods of weighted residuals such as Galerkin and collocation methods.

Previous applications of the method of weighted residuals to combustion-instability problems have dealt with pressure-sensitive combustion. In this work this method will be extended to handle situations in which velocity sensitivity is important. Both the collocation and Galerkin methods will be considered. Stability boundaries and pressure wave forms will be computed numerically for transverse motion in a cylindrical combustion chamber. In addition a finite-difference method will be used to solve the one-dimensional problem of transverse motion in a thin annular chamber. These results will be compared with those obtained by a method of weighted residuals solution of the same problem in order to assess the accuracy of the approximate solution.

In this study, attempts to solve the velocitysensitive combustion instability problem by various
numerical methods are considered. The governing equations
that describe the flow conditions inside liquid-propellant
rocket motors will be derived in Chapter Two. An analytical
solution is obtained in Chapter Three for nonlinear acoustic
motion in a cylindrical chamber. The collocation method is

applied to combustion instability in a cylindrical chamber in Chapter Four. In Chapter Five, the Galerkin method is applied to the same problem. In Chapter Six, attention is given to the analysis of combustion instability in a thin annulus. This allows the comparison of the Galerkin method and a finite-difference solution in a one-dimensioned context. Finally, a summary of this research is contained in Chapter Seven.

Chapter 2

COMBUSTOR EQUATIONS

It is the objective of the discussion presented in this chapter to analyze the combustor conservation equations and reduce them to a tractable system. The simplification will be done in such a manner that will allow the resulting equations to retain both the mathematical and physical essence of the original problem.

Under the assumptions that the usual balance principles of mechanics can be applied separately to each phase, the following equations are derived by applying the laws of conservation of mass, momentum, and energy to an arbitrary stationary control volume.

Conservation of mass of the fluid requires that the time rate of change of mass in a volume v equals the rate at which mass enters v plus the rate at which mass is generated inside v and can be expressed as follows (for a fixed volume):

$$\partial_t * \int_v^* \vec{\rho} dv = - \int_s^* \vec{\rho} \vec{u} \cdot \vec{n} ds + \int_v^* \vec{w} dv$$

where \vec{n} is an outward unit vector normal to s, $\vec{\rho}$ is the density of fluid, \vec{u} is the velocity of fluid, \vec{w} is the rate at which mass is generated per unit volume, and \vec{t} is the time.

Using the divergence theorm and the arbitrariness of the stationary control volume, yields

Similarly, the balance law of mass for fuel phase is

$$\partial_{t} * \rho_{L}^{*} + \vec{\nabla} \cdot (\rho_{L}^{*} \vec{u}_{L}) = - \vec{w}$$
 (2.2)

where $\overset{\boldsymbol{\star}}{\rho_L}$ is the density of fuel and $\overset{\boldsymbol{\star}}{u_L}$ is the velocity of fuel.

The balance law of linear momentum states that the rate of change of linear momentum in the volume v equals the rate of entry of linear momentum across the surface s plus the sum of all external forces acting on the volume v and can be expressed as

$$\partial_{t} * \int_{V} \dot{\vec{v}} \dot{\vec{u}} dv = - \int_{S} \dot{\vec{v}} \dot{\vec{u}} \cdot \dot{\vec{n}} ds + \int_{V} \dot{\vec{F}} dv + \int_{V} \dot{\vec{w}} \dot{\vec{u}}_{L} dv - \int_{S} \dot{\vec{P}} \dot{\vec{n}} ds$$

where $\overset{*}{F}$ is the force per unit volume applied to the gas by the fuel and $\overset{*}{P}$ is the pressure of the gas.

Using the same argument, the equation becomes

$$\partial_{t,*}(\mathring{\rho}\mathring{u}) + \mathring{\nabla} \cdot (\mathring{\rho}\mathring{u}\mathring{u}) = \mathring{F} + \mathring{w}\mathring{u}_{T} - \mathring{\nabla}P.$$
 (2.3)

Similarly, the balance of momentum for the fuel phase is

$$\partial_{t} * (\overset{*}{\rho}_{L} \overset{*}{u}_{L}) + \overset{*}{\nabla} \cdot (\overset{*}{\rho}_{L} \overset{*}{u}_{L} \overset{*}{u}_{L}) = -\overset{*}{F} - \overset{*}{w} \overset{*}{u}_{L}.$$
 (2.4)

Substituting (2.1) in (2.3) yields

Similarly for the fuel phase

The law of conservation of energy states that the rate of change of energy in a volume v equals the rate at which energy enters the volume v plus the rate at which energy is generated internally plus the rate at which work is done by external forces and gives

$$\partial_{t} * \int_{v}^{*} (e^{+\frac{\pi}{2}u^{2}}) dv = -\int_{s}^{*} (e^{+\frac{\pi}{2}u^{2}}) (u^{+} \cdot n^{+}) ds - \int_{s}^{*} (Pn^{+}) \cdot u^{+} ds$$

$$+ \int_{V}^{*} \vec{F} \cdot \vec{u}_{L} dv + \int_{V}^{*} (e_{L}^{*} + \frac{1}{2}u_{L}^{2}) dv + \int_{V}^{*} Qdv$$

where $\overset{*}{e}$ + $\frac{*}{2}\overset{*}{u}^2$ is the energy of gas per unit volume, $\overset{*}{e}_L$ + $\frac{*}{2}\overset{*}{u}_L^2$ is the energy of fuel per unit volume, and $\overset{*}{Q}$ is the heat transfer rate from the fuel to the gas.

Following the same argument, the equation becomes

$$\partial_{t}*(\hat{p}(e^{+} + b_{2}u^{2})) + \nabla \cdot (\hat{p}(e^{+} + b_{2}u^{2})u^{2}) = -\nabla \cdot (Pu)$$

$$+ \dot{F} \cdot \dot{u}_{L} + \ddot{w} (\dot{e}_{L} + \dot{z}\dot{u}_{L}^{2}) + \ddot{Q}.$$
 (2.6)

Similarly, the balance of energy for the fuel phase becomes

$$\frac{1}{2} \frac{(\mathring{\rho}_{L})^{2}}{(\mathring{\rho}_{L})^{2}} = - \frac{\mathring{F} \cdot \mathring{u}_{L}^{2}}{\mathring{r}_{L}} - \mathring{w}(\mathring{e}_{L}^{2} + \frac{1}{2}\mathring{u}_{L}^{2}) - \mathring{Q}.$$

$$\frac{\mathring{\pi}}{(\mathring{\rho}_{L})^{2}} = - \mathring{F} \cdot \mathring{u}_{L}^{2} - \mathring{w}(\mathring{e}_{L}^{2} + \frac{1}{2}\mathring{u}_{L}^{2}) - \mathring{Q}.$$
(2.7)

Substituting (2.1), (2.2) into (2.6), (2.7) respectively yields

$$\mathring{\rho}_{L}(\partial_{t}*(\mathring{e}_{L} + \frac{1}{2}\mathring{u}_{L}^{2}) + \mathring{u}_{L}^{*} \cdot \mathring{\nabla}(\mathring{e}_{L}^{*})$$

$$+ \frac{*}{2}u_{L}^{2})) = - \vec{F} \cdot \vec{u}_{L} - \vec{Q}.$$
 (2.9)

It is more convenient to work with the thermal energy equation for each phase. For gas phase this is done by

dotting \vec{u} into (2.3) to obtain the mechanical energy equation and subtracting this from (2.8) and using (2.1) to get

$$+ (\dot{u}_{L}^{2} - \dot{u}^{2})/2 + \dot{\dot{u}} \cdot (\dot{\dot{u}} - \dot{\dot{u}}_{L}) - P/\rho). \qquad (2.10)$$

In a similar way, it can be shown that

where D /Dt* = ∂_{t*} + $\overrightarrow{u} \cdot \overrightarrow{\nabla}$, D_L /Dt* = ∂_{t*} + $\overrightarrow{u}_{L} \cdot \overrightarrow{\nabla}$ are the comoving derivatives for each phase.

For simplicity, gas phase viscosity and heat conduction were neglected in the previous analysis. Since these produce dissipation, equations derived in this way should give a conservative estimate of the stability of the system.

It appears (see, for instance, Powell (1)) that the primary effect of the burning fuel is that of interphase mass transfer. Thus, the balance laws for mass, momentum, and thermal energy for each phase will be further simplified by neglecting all interphase transfer terms other than those appearing in the mass-balance equations.

This leads to the system of equations:

$$D_{\rho}^{*}/Dt^{*} + \rho^{*} \vec{\nabla} \cdot \vec{u} = \vec{w}$$

$$D_{\rho}^{*}/Dt^{*} + \rho^{*} \vec{\nabla} \cdot \vec{u}_{L} = -\vec{w}$$

$$\rho^{*}/D_{L}/Dt^{*} + \rho^{*}/D_{L} \vec{v}_{L}/Dt^{*} = 0$$

$$\rho^{*}/D_{L} \vec{u}_{L}/Dt^{*} = 0$$

where the constitutive equations $\overset{*}{e} = C_v^{\overset{*}{T}}$ and $\overset{*}{e}_L = C_{vL}^{\overset{*}{T}}_L$ have been used (with $\overset{*}{T}$ denoting temperature and C_v specific heat). The equation of state for an ideal gas is

* * * * P =
$$\rho RT$$
 (2.13)

where R is the gas constant.

The governing equations will now be nondimensionalized with respect to a steady reference state, which will be denoted by the subscript "r". All lengths will be referred to some characteristic length $L_{\rm r}$, such as the chamber radius. The characteristic velocity is the speed of sound at the

reference state, and the characteristic time is the wave travel time $L_{\rm r}/c_{\rm r}$. The dimensionless quantities are defined below:

$$\frac{*}{\vec{V}} = \vec{V} / L_{r} \qquad \qquad \hat{t} = L_{r} t / C_{r}$$

$$\frac{*}{\vec{U}} = C_{r} \vec{U} \qquad \qquad \hat{u}_{L} = C_{r} \vec{u}_{L}$$

$$\frac{*}{\rho} = \rho_{r} \rho \qquad \qquad \hat{\rho}_{L} = \rho_{r} \rho_{L}$$

$$\frac{*}{\rho} = P_{r} P \qquad \qquad \hat{v} = \rho_{r} C_{r} w / L_{r}$$

$$\frac{*}{T} = C_{r}^{2} T / (\gamma R) \qquad \qquad T_{L} = C_{r}^{2} T_{L} / (\gamma R)$$

$$C_{r}^{2} = \gamma P_{r} / \rho_{r} \qquad \qquad P_{r} = \rho_{r} C_{r}^{2} / \gamma .$$

The dimensionless conservation equations become

Continuity

$$D\rho/Dt + \rho \vec{\nabla} \cdot \vec{u} = w$$

$$D\rho_L/Dt + \rho_L \vec{\nabla} \cdot \vec{u}_L = -w \qquad (2.14)$$

Momentum

$$D\vec{u}/Dt = - \vec{\nabla}P/\gamma$$

$$D\vec{u}_{T}/Dt = 0$$
 (2.15)

Energy

$$DT/Dt = (\gamma - 1)P(D/Dt)(log\rho)$$

$$DT_{\mathsf{T}}/Dt = 0 \tag{2.16}$$

Equation of state

$$P = \rho T . \tag{2.17}$$

By solving (2.16a) one obtains

$$T = \rho^{\gamma - 1} . \tag{2.18}$$

Equation (2.17) then becomes

$$P = \rho^{\gamma}. \tag{2.19}$$

Substituting (2.19) to (2.15a) produces

$$\overrightarrow{Du}/Dt = -\overrightarrow{\nabla}(\rho^{\gamma-1})/(\gamma-1). \qquad (2.20)$$

Taking the curl on both sides of equation (2.20) gives

$$\vec{\nabla} \times \vec{Du}/Dt = 0. \tag{2.21}$$

It can be shown using vector identity that (2.21) leads to the following equation which describes the generation of the vorticity $\vec{\Omega} = \vec{\nabla} \ X \ \vec{u}$ in the flow.

$$D\vec{\Omega}/Dt = \vec{\Omega} \cdot (\vec{\nabla} \vec{u}) - \vec{\Omega}(\vec{\nabla} \cdot \vec{u}) . \qquad (2.22)$$

This equation describes the variation of vorticity for a given fluid particle. Suppose that at t = 0, this particle has zero vorticity and at the point where it is located the velocity gradients are bounded. Then by (2.22) the rate of change of vorticity of the particle vanishes. It follows from this that the vorticity will be zero at the next instant. By induction, it is seen that as long as the velocity gradients are bounded at each point occupied by the particle, its vorticity will remain zero for all time. all fluid particles in the system have zero vorticity at t = 0, the vorticity will vanish at all points in the flow field and for all $t \geq 0$, therefore, as long as the velocity gradients are bounded through the flow field (as they would not be, for example, at a shock wave). Assuming this condition to be satisfied implies irrotational flow $(\stackrel{\rightarrow}{\Omega} = 0)$ and allows the definition of a velocity potential ϕ defined such that

$$\vec{u} = \vec{\nabla} \phi . \qquad (2.23)$$

Substituting (2.23) to (2.20) gives

$$\partial_{t} \phi + \frac{1}{2} \vec{\nabla} \phi \cdot \vec{\nabla} \phi + \rho^{\gamma-1}/(\gamma-1) = F(t)$$
.

Since the velocity is the space derivative of ϕ , it is permissible to add to ϕ any arbitrary function of time. This is equivalent to a statement that F(t) is arbitrary. For future convenience F(t) was selected to be $1/(\gamma-1)$. This choice yields

$$\rho^{\gamma-1} = 1 - (\gamma-1)(\partial_{\tau}\phi + \frac{1}{2}\vec{\nabla}\phi \cdot \vec{\nabla}\phi) . \qquad (2.24)$$

Substituting (2.24) into (2.14a), (2.18) and (2.19) gives

$$\begin{array}{lll} \partial_{tt}\phi & - \nabla^2\phi(1-(\gamma-1)(\partial_t\phi + \frac{1}{2}\vec{\nabla}\phi\cdot\vec{\nabla}\phi)) + \vec{\nabla}\phi\cdot\vec{\nabla}(2\partial_t\phi \\ \\ & + \frac{1}{2}\vec{\nabla}\phi\cdot\vec{\nabla}\phi) + (1-(\gamma-1)(\partial_t\phi + \frac{1}{2}\vec{\nabla}\phi\cdot\vec{\nabla}\phi)) & (\gamma-2)/(\gamma-1)_{W=0} \end{array}$$

$$T = 1-(\gamma-1)(\partial_t\phi + \frac{1}{2}\vec{\nabla}\phi\cdot\vec{\nabla}\phi)$$

$$P = (1-(\gamma-1)(\partial_t\phi + \frac{1}{2}\vec{\nabla}\phi\cdot\vec{\nabla}\phi))^{\gamma/(\gamma-1)}. \qquad (2.25)$$

This system of equations contains nonlinear gas dynamics terms of all orders.

Due to their highly nonlinear and mathematically complicated nature, the system of equations obtained above cannot be solved exactly. In order to obtain simpler, but approximate, equations which can be more easily analyzed, all nonlinear terms of order higher than two are neglected. This has the effect of retaining the most important nonlinear effect, while greatly simplifying the form of the equations. The system of equations then becomes

$$\rho = 1 - (\partial_{t}\phi + \frac{1}{2}\vec{\nabla}\phi \cdot \vec{\nabla}\phi) + \frac{1}{2}(2-\gamma)(\partial_{t}\phi)^{2}$$

$$T = 1 - (\gamma-1)(\partial_{+}\phi + \frac{1}{2}\vec{\nabla}\phi \cdot \vec{\nabla}\phi)$$

$$P = 1 - \gamma(\partial_{t}\phi + \frac{1}{2}\vec{\nabla}\phi \cdot \vec{\nabla}\phi) + \frac{1}{2}\gamma(\partial_{t}\phi)^{2}$$

$$\partial_{tt}\phi - \nabla^{2}\phi (1 - (\gamma - 1)\partial_{t}\phi) + 2\vec{\nabla}\phi \cdot \vec{\nabla}(\partial_{t}\phi)$$

$$+ (1 - (\gamma - 2)\partial_{t}\phi)w = 0. \qquad (2.26)$$

For a cylindrical combustor, it is desired to investigate the stability of the steady-state solution of (2.26d). To do this the steady-state solution must be obtained. It is assumed that the steady-state burning rate depends on z only ($w = \underline{w}(z)$) and that the flow is axial ($\phi = \phi(z)$). Thus (2.26) simplifies to

$$d\underline{u}/dz = \underline{w} \qquad (\underline{u} = d\underline{\phi}/dz)$$

$$\underline{T} = 1 - \frac{1}{2}(\gamma - 1)\underline{u}^{2}$$

$$\underline{P} = 1 - \frac{1}{2}\gamma\underline{u}^{2}$$

$$\varrho = 1 - \frac{1}{2}\underline{u}^{2}. \qquad (2.27)$$

It can be seen from (2.27) that the deviations of the steady-state solution from a uniform state are $O(\underline{u}^2)$. To allow a discussion of orders of magnitude of various terms define the ordering parmeter ε such that

$$\varepsilon = \max(\underline{u}).$$
 (2.28)

This implies that

$$\underline{\mathbf{u}} = O(\varepsilon), \quad \underline{\mathbf{w}} = O(\varepsilon), \quad \underline{\phi} = O(\varepsilon).$$
 (2.29)

The stability analysis will now be carried out by assuming that

$$\phi = \underline{\phi}(z) + \varepsilon \phi'(x, y, z, t)$$

$$w = \underline{w}(z) + \varepsilon^{2} w'(x, y, z, t) \qquad (2.30)$$

where
$$\phi' = O(1)$$
, $w' = O(1)$. (2.31)

It is being assumed that the unsteady perturbation of the velocity potential from the steady state is of the same order of magnitude as the deviation of the steady state from a uniform state and that the unsteady burning rate perturbation is of the same order as the unsteady nonlinear gas-dynamic terms. The first assumption is necessary to be consistent with the quadratic approximation inherent in (2.26) while the second eliminates nonlinear terms involving products of w' and o'. These assumptions are made for simplicity and are not meant to imply that other orders of magnitude for the various terms are impossible or even unlikely. The objective of this work is to pick one case which is mathematically tractable and subject it to extensive analysis. While it is felt that most of the features of the stability problem will be reflected by this case, it is recognized that many other possibilities remain to be explored.

Substituting (2.30) and

$$\rho = \rho + \epsilon \rho', \quad P = P + \epsilon P', \quad T = T + \epsilon T'$$
 (2.32)

 $\{\rho'=0(1),\ P'=0(1),\ T'=0(1)\}$ into (2.26),retaining all terms of $0(\epsilon^3)$ or lower,and dividing all resulting equations by ϵ leads to

$$\partial_{tt}\phi' - \nabla^{2}\phi' + 2\underline{u}\partial_{zt}\phi' + \underline{w}\partial_{t}\phi'$$

$$+ \varepsilon\{(\gamma-1)\nabla^{2}\phi'\partial_{t}\phi' + 2\overline{\nabla}\phi'\cdot\overline{\nabla}(\partial_{t}\phi') + w'\} = 0$$

$$\rho' = - \{\partial_{t}\phi' + \underline{u}\partial_{z}\phi' + \frac{1}{2}\varepsilon(\overline{\nabla}\phi'\cdot\overline{\nabla}\phi' - (2-\gamma)(\partial_{t}\phi')^{2})\}$$

$$P' = - \gamma\{\partial_{t}\phi' + \underline{u}\partial_{z}\phi' + \frac{1}{2}\varepsilon(\overline{\nabla}\phi'\cdot\overline{\nabla}\phi' - (\partial_{t}\phi')^{2})\}$$

$$T' = -(\gamma-1)(\partial_{t}\phi' + \underline{u}\partial_{z}\phi' + \frac{1}{2}\varepsilon\nabla\phi'\cdot\nabla\phi'). \qquad (2.33)$$

The equations derived from this perturbation analysis are expected to be valid as long as the amplitudes of the perturbation quantities are finite and smaller than unity.

For simplicity the primes appearing in the above equations will be dropped and it will be understood that unprimed quantities are associated with perturbation from the steady state.

Chapter 3

ANALYTICAL SOLUTION OF WAVE EQUATION

In this chapter a second-order solution for the velocity potential associated with transverse acoustic wave motion of an ideal gas in a circular cylinder is obtained. The solution is found in the form of a Fourier-Bessel series. This provides a standard to compare the proposed methods which will be discussed in the following chapters.

The governing equation is obtained from (2.33a) by assumption that the burning rate function is absent. Then the equation reduces to the second-order form of the one discussed by Maslen and Moore (2) for pure gas motion in a cylinder.

$$\partial_{t,t}\phi - \nabla^2\phi + \varepsilon(2\vec{\nabla}\phi\cdot\vec{\nabla}\partial_t\phi + (\gamma-1)\nabla^2\phi\partial_t\phi) = 0 \qquad (3.1)$$

For a cylinder of radius L_r , a set of cylindrical polar coordinates (r,θ,z) with the z axis being coincident with the cylinder's axis of symmetry is used. In seeking a second-order transverse wave solution, one assumes

$$\phi = \phi_1(r,\theta,t) + \epsilon \phi_2(r,\theta,t) + \dots \qquad (3.2)$$

Substituting (3.2) into (3.1) and (2.33) leads to

$$\partial_{tt}\phi_{1} - \nabla^{2}\phi_{1} = 0$$

$$\partial_{tt}\phi_{2} - \nabla^{2}\phi_{2} = -(\partial_{t}(u_{1}^{2}) + (\gamma-1)\nabla^{2}\phi_{1}\partial_{t}\phi_{1})$$
 (3.3)

$$P = 1 + \epsilon P_1 + \epsilon^2 P_2 \tag{3.4}$$

where
$$P_1 = -\gamma \partial_t \phi_1$$
, $P_2 = -\gamma (\partial_t \phi_2 + \frac{1}{2} (u_1^2 - (\partial_t \phi_1)^2))$ (3.5)

$$\nabla^2 \phi_i = \partial_{rr} \phi_i + \partial_r \phi_i / r + \partial_{\theta\theta} \phi_i / r^2$$
, $i = 1, 2$

$$u_1^2 = (\partial_r \phi_1)^2 + (\partial_\theta \phi_1/r)^2.$$
 (3.6)

Three different cases are discussed below. For the initial conditions

$$\phi(r,\theta,0) = J_1(S_1r)\cos\theta, \ \theta_t\phi(r,\theta,0) = 0 \tag{3.7}$$

the solution of (3.3a) is

$$\phi_1 = J_1(S_1r)\cos\theta\cos(S_1t).$$

Substituting this equation to (3.3b) produces

$$\partial_{tt} \phi_{2}^{-\nabla^{2}} \phi_{2}^{2} = (b_{00}^{+} + \sum_{m=0}^{\infty} \sum_{n=1}^{\infty} J_{m}(S_{mn}r) \cos(m\theta)) \sin(2S_{11}t)$$
 (3.8)

where J_n is used to denote an n'th order Bessel function of the first kind, S_{mn} is used to denote the m'th root of the equation $J_n'=0$, and a prime indicates differentiation

with respect to the argument of the Bessel function. The coefficients \mathbf{b}_{mn} are integrals of Bessel functions which must be computed numerically. The details of this calcuation are discussed in appendix A.

Solving (3.8) one obtains

$$\phi_2 = (b_0 / (4S^2))(2S_1 t - sin(2S_1 t))$$

$$\begin{array}{l} + \sum\limits_{m=0}^{2} \sum\limits_{n=1}^{\infty} (b_{mn}/(S_{mn}^{2} - 4S_{11}^{2})) J_{m}(S_{mn}r) \cos(m\theta) (\sin(2S_{11}t) \\ m \neq 1 \end{array}$$

-
$$(2S_{11}/S_{mn})sin(S_{mn}t)$$
. (3.9)

The other two solutions to be discussed subsequently are found in the same way. For the initial conditions

$$\phi(r,\theta,0) = 0, \ \partial_t \phi(r,\theta,0) = S_{11} J_1(S_{11} r) \cos\theta$$
 (3.10)

it is found that

$$\phi_1 = J_1(S_{11})\cos\theta\sin(S_{11})$$
 (3.11)

and that ϕ_2 is given by the negative of (3.9). For the initial conditions

$$\phi(r,\theta,0) = J_1(S_1r)\cos\theta,$$

$$\partial_{t} \phi(r, \theta, 0) = S J (S_{11} r) \sin \theta \qquad (3.12)$$

the solution is

$$\phi_1 = J_1(S_{11}r)\cos(S_{11}t - \theta)$$

and
$$\phi_2 = \sum_{n=1}^{\infty} \{2b_{2n}/(S_{2n}^2 - 4S_{11}^2)\}J_2(S_{2n}r)(\sin(2\theta)\{\cos(2S_{11}t)\}$$

$$-\cos(S_{2n}t)\} + \cos(2\theta)\{\sin(2S_{11}t) - (2S_{11}/S_{2n})\sin(S_{2n}t)\}).$$

It can be seen that for these initial conditions ϕ_1 represents a spinning wave but $\phi_1 + \epsilon \phi_2$ does not. Some typical results for the pressure functions P_1 and P_2 were computed using the solutions for initial conditions (3.7) (labeled standing wave) and initial conditions (3.12) (labeled traveling wave) for r=1, $\theta=0$, and $\gamma=1.2$. These data are presented graphically in Figure 1. For these computations the series expansions were terminated at n=5. There is virtually no difference between the results for n=4 and n=5, and even the results for n=1 provide a reasonably accurate solution.

The results discussed above were used to check the numerical calculations to be discussed in the next two chapters. These results generalize those of Maslen and Moore (2) to initial conditions which do not lead to periodic solutions.

Chapter 4

COLLOCATION

In previous investigations of velocity-sensitive combustion instability {see, for instance, (6) and the references therein} the most widely used burning-rate equation is that associated with the vaporization limited combustion model discussed by Priem and Guentert (3). In the notation of the present thesis the second-order version of this equation can be expressed as

$$W = (\underline{W}/\epsilon^2)((1 - \frac{1}{2}\epsilon \partial_t \phi)((1 - u_z/u_L)^2 + (u_t/u_L)^2)^{\frac{1}{4}} - 1) (4.1)$$

where $\mathbf{u}_{\mathbf{z}}$ is the component of perturbed gas velocity in the axial direction and $\mathbf{u}_{\mathbf{t}}$ is the perturbed component normal to the axis of symmetry, and $\mathbf{u}_{\mathbf{L}}$ is the magnitude of the droplet velocity vector which assumed to be axial. Substituting (4.1) into (2.33a) yields

$$\begin{array}{l} \partial_{tt}\phi + \underline{w}\,\partial_{t}\phi + 2\underline{u}\,\partial_{zt}\phi - \nabla^{2}\phi(1-\varepsilon(\gamma-1)\,\partial_{t}\phi) \\ \\ + \,2\,\varepsilon\vec{\nabla}\phi\cdot\vec{\nabla}\,\partial_{t}\phi + (\underline{w}/\varepsilon)((1-\frac{1}{2}\varepsilon\,\partial_{t}\phi)((1-\partial_{z}\phi/u_{L})^{2} \\ \\ + \,(\vec{\nabla}\phi\cdot\vec{\nabla}\phi - (\partial_{z}\phi)^{2})/u_{L}^{2})^{\frac{1}{4}} - \,1) = 0\,. \end{array} \tag{4.2}$$

No exact solutions are to be expected to (4.2). Fully numerical solutions, on the other hand, are expected to be quite time consuming for multidimensional problems. these reasons it was decided to attempt approximate partially analytical solutions using the method of weighted residuals (see for instance, (6)). For problems of pressure sensitive combustion instability Powell (1) successfully employed the Galerkin method. This method is limited in applicability, however, to equations containing nonlinearities involving only polynomial functions of the dependent variable and its derivatives. It is clear that (4.2) is not of this form. The orthogonal collocation method was, therefore, chosen because of its simplicity and adaptability to equations containing nonlinearities of any algebraic The application of this method to the problem of transverse wave motion in a cylindrical chamber will be discussed below.

Consider the transverse wave motion in a cylindrical combustor. It is convenient to describe this problem in terms of cylindrical polar coordinates (r, θ, z) . Then

$$\phi = \phi(r, \theta, t). \tag{4.3}$$

It will also be assumed that \underline{w} and u_L are constants. Then (3.2) becomes

$$\partial_{tt}\phi + \underline{w}\partial_{t}\phi - (\partial_{rr}\phi + \partial_{r}\phi/r + \partial_{\theta\theta}\phi/r^{2})(1 - \varepsilon(\gamma-1)\partial_{t}\phi)$$

$$+2\varepsilon(\partial_{\mathbf{r}}\phi\partial_{\mathbf{r}t}\phi+\partial_{\theta}\phi\partial_{\theta t}\phi/\mathbf{r}^{2})+(\underline{\mathbf{w}}/\varepsilon)((1-\frac{1}{2}\varepsilon\partial_{t}\phi)((1-\frac{1}{2}\varepsilon\partial_{t}\phi))((1-\frac{1}{2}\varepsilon\partial_{t}\phi))$$

+
$$((\partial_r \phi)^2 + (\partial_\theta \phi/r)^2)/u_L^2)^{\frac{1}{4}} - 1) = 0$$
. (4.4)

Now, assume an approximate solution of the form

$$\phi = \sum_{m=0}^{P} \sum_{n=1}^{Q} J_m(S_{mn}r) \cos(m\theta) f_{mn}(t). \tag{4.5}$$

Equation (4.5) is a superposition of the normal modes associated with the corresponding linear acoustic problem. A solution of this form allows one to investigate the influence of nonlinearities on the behavior of modal amplitudes which would exhibit simple harmonic motion in the linear case. Only standing modes can be described by (4.5). To represent spinning modes another series involving $\sin(m_{\theta})$ must be added. This is omitted for simplicity in this discussion. It is convenient to rewrite (4.5) as

$$\phi = \int_{j=1}^{N} J_{m_{j}}(S_{m_{j}n_{j}})\cos(m_{j}\theta)f_{j}(t)$$
 (4.6)

where N = (P + 1)Q.

The equation is now evaluated at N points (the collocation points) to yield

$$\phi_{i} = \sum_{j=1}^{N} C_{i,j} f_{j}$$
 $i = 1, 2, ... N$ (4.7)

where
$$C_{ij} = J_{m_j} (S_{m_j n_j} r_i) cos(m_j \theta_i).$$
 (4.8)

Next, (4.7) is inverted to obtain

$$\mathbf{f}_{\mathbf{i}} = \sum_{\mathbf{j}=1}^{N} \mathbf{C}_{\mathbf{i}\mathbf{j}}^{-1} \phi_{\mathbf{j}}. \tag{4.9}$$

Then the various derivatives of (4.6) can be expressed in terms of value of $\phi_{_{\mbox{\scriptsize i}}}$ as

$$(\partial_{\mathbf{r}} \phi)_{\dot{\mathbf{i}}} = \sum_{\mathbf{j}=1}^{N} C_{\dot{\mathbf{i}}\dot{\mathbf{j}}}^{\mathbf{r}} \phi_{\dot{\mathbf{j}}}, \quad (\partial_{\mathbf{r}\mathbf{r}} \phi)_{\dot{\mathbf{i}}} = \sum_{\mathbf{j}=1}^{N} C_{\dot{\mathbf{i}}\dot{\mathbf{j}}}^{\mathbf{r}\mathbf{r}} \phi_{\dot{\mathbf{j}}}$$

$$(\partial_{\theta} \phi)_{\dot{\mathbf{i}}} = \sum_{\mathbf{j}=1}^{N} C_{\dot{\mathbf{i}}\dot{\mathbf{j}}}^{\theta} \phi_{\dot{\mathbf{j}}}, \quad (\partial_{\theta} \phi)_{\dot{\mathbf{i}}} = \sum_{\mathbf{j}=1}^{N} C_{\dot{\mathbf{i}}\dot{\mathbf{j}}}^{\theta} \phi_{\dot{\mathbf{j}}} \quad \dot{\mathbf{i}} = 1, \dots N$$

$$(4.10)$$

where
$$C_{ij}^{r} = (\partial_{r_{ik}}^{c})C_{kj}^{-1}, C_{ij}^{rr} = (\partial_{r_{i}r_{i}}^{c}C_{ik})C_{kj}^{-1}$$

$$C_{ij}^{\theta} = (\partial_{\theta}C_{ik})C_{kj}^{-1}, C_{ij}^{\theta\theta} = (\partial_{\theta_{i}\theta_{i}}^{c}C_{ik})C_{kj}^{-1}. \tag{4.11}$$

Substituting into (4.4), a set of N ordinary second-order differential equations having the following form is generated.

$$\frac{\partial}{\partial t} + \underline{w} \dot{\phi}_{i} - \frac{\partial}{\partial z_{i}} C_{ij}^{\nabla} \dot{\phi}_{j} (1 - \varepsilon (\gamma - 1) \dot{\phi}_{j}) + 2\varepsilon \sum_{j=1}^{N} \sum_{k=1}^{N} C_{ijk} \dot{\phi}_{j} \dot{\phi}_{k}$$

$$+ (\underline{w}/\varepsilon) ((1 - \frac{1}{2}\varepsilon \dot{\phi}_{i}) (1 + \sum_{j=1}^{N} \sum_{k=1}^{N} C_{ijk} \dot{\phi}_{j} \dot{\phi}_{k} / u_{L}^{2})^{\frac{1}{4}} - 1) = 0 \qquad (4.12)$$

where $C_{ij}^{\nabla} = C_{ij}^{rr} + C_{ij}^{r}/r_i + C_{ij}^{\theta\theta}/r_i^2$

$$C_{i,j,k} = C_{i,j}^{r} C_{i,k}^{r} + C_{i,j}^{\theta} C_{i,k}^{\theta} / r_{i}^{2}$$
 (4.13)

Solutions were obtained by a fourth-order Runge-Kutta method to the set of equations (4.12). It was found that the results were very sensitive to the number and locations of collocation points, especially to the radial distribution of points. It was found that even in cases when \underline{w} was equated to zero (no combustion) it was possible to find many choices of collocation points which lead to the computed modal amplitudes increasing without bound.

In order to illustrate the difficulties involved in a simple case, consider a one-dimensional problem of longitudinal wave motion with $\gamma=1$, $\underline{u}=0$ and $\underline{w}=0$. In this case (4.2) reduces to

$$\partial_{tt} \phi - \partial_{zz} \phi + 2\varepsilon (\partial_{z} \phi \partial_{zt} \phi) = 0 . \qquad (4.14)$$

First consider the boundary condition

$$\partial_{z} \phi(0,t) = 0, \qquad \phi(\pi,t) = 0.$$
 (4.15)

A one-term solution satisfying (4.15) is

$$\phi = f(t)\cos(\frac{1}{2}z).$$
 (4.16)

Applying the collocation method with a collocation point z_0 produces an equation for $\phi_0 = \phi(z_0, t)$ of the form

$$\ddot{\phi}_{0} + \chi_{\phi}_{0} + \chi(\tan^{2}(z_{0}/2)\phi_{0}\dot{\phi}_{0}) = 0. \tag{4.17}$$

The Galerkin method (to be discussed in detail in the next chapter) when applied to the same problem yields an equation for f of the form

$$f + \frac{1}{4}f + 4 ff/(3\pi) = 0. (4.18)$$

As another example consider the boundary conditions

$$\partial_{z}\phi(0,t) = 0, \qquad \partial_{z}\phi(\pi,t) = 0.$$
 (4.19)

A solution satisfying (4.19) is

$$\phi = f(t)\cos(z) \tag{4.20}$$

for which the collocation method leads to

and the Galerkin method leads to

$$f + f = 0. (4.22)$$

From the above two cases, one can observe that the last term in each of the equations obtained by collocation can take on any value between 0 and ∞ depending on the location z_0 of the collocation point. No such difficulty is encountered when using the Galerkin method. Even if more points were used, the same behavior was obtained. Therefore

there appears to be no way to rationally decide which sets of collocation points can produce the correct result. For this reason, after the expenditure of much effort, the collocation method was abandoned and the Galerkin method was adopted. This will be discussed in the next chapter.

Chapter 5

THE GALERKIN METHOD

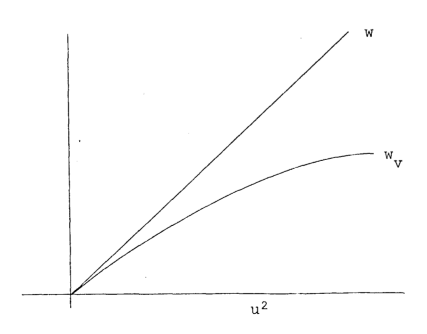
The Galerkin method is a special application of the method of weighted residuals (usually referred to as MWR). It has been extensively used in the solution of various stability and aeroelasticity problems {see for instance Powell (1) and proved itself as a useful tool for the solution of both linear and nonlinear problems. Although it is an approximate mathematical technique, it has nevertheless produced results which were in excellent agreement with available exact solutions. These approximate solutions are usually simpler in form than the exact solutions obtained by numerical integration, and their quantitative evaluation requires considerably less computation time. However, this method is known to be reliable and applied conveniently only to equations involving nonlinearities of a polynomial type. Therefore, if this method is used in conjunction with the varporizationlimited burning rate function of (4.1) the problem becomes intractable. Thus, the following simpler purely phenomenological burning-rate function was employed for the case of instantaneous combustion response.

$$w = \underline{w}nu^2 \tag{5.1}$$

where n is a constant which will subsequently be referred to as the interaction coefficient. It can be determined either by comparison of predictions of the theory directly with experiment or by comparison with burning rate laws meant to apply to special types of combustion processes. As an example of the latter method consider the vaporization—limited burning rate law (4.1). As it stands (4.1) exhibits both pressure and velocity sensitivity. A purely velocity sensitive law can be obtained by assuming ϵ << 1 to get

$$W_{V} = (\underline{w}/\varepsilon^{2})((1 + (u/u_{L})^{2})^{\frac{1}{4}} - 1)$$
 (5.2)

where the v is used to denote the vaporization model. If one equates the slopes of (5.1) and (5.2) at $u^2 = 0$ and plots both functions, the graph would look as follows.



It can be seen that (5.1) would overestimate the burning rate. Thus the upper bound for n can be computed. From (5.1) and (5.2)

$$d_{u^2}w = \underline{w}n, \quad d_{u^2}w_v = (\underline{w}/(4\epsilon^2u_L^2))/(1+(u/u_L)^2)^{3/4}.$$

Thus
$$d_{u2}w(0) = \underline{w}n$$
, $d_{u2}w_{v}(0) = \underline{w}/(4\epsilon^{2}u_{L}^{2})$.

Equating these results, one gets

$$n_{\text{max}} = 1/(4 \epsilon^2 u_L^2).$$
 (5.3)

The phenomonological law (5.1) can be related to other special burning-rate laws in a similar manner.

If it is desired to consider history-dependent combustion processes, this can be done through the general formula

$$w = n \int_{0}^{t} G(t-\xi) d_{\xi}(u^{2}) d\xi$$
 (5.4)

where G is memory function and ξ is a dummy variable. If G(t) = H(t) (H being the unit step function) all increments of change in u^2 occurring in the past are counted equally and (5.1) is recovered. If $G(t) = H(t) - H(t-\tau)$ all increments of change in u^2 are counted equally up to τ units of time in the past while those previous to that time are not counted at all. Substituting this expression into (5.4)

and integrating one obtains

$$w = \underline{w}n(u^2 - u_T^2)$$
 (5.5)

where $u_{\tau}=u(t-\tau)$ and τ is called the time delay. Equation (5.5) will be used to account for the history of the burning process in a rough way. More sophisticated treatments of this phenomenon could be obtained by employing a more realistic function for G(t). Substituting (5.5) and (5.1) into (2.33a) one obtains

$$\partial_{tt}\phi - \nabla^{2}\phi + 2\underline{u}\partial_{zt}\phi + \underline{w}\partial_{t}\phi + \varepsilon((\gamma-1)\nabla^{2}\phi\partial_{t}\phi + 2\overline{\nabla}\phi\cdot\overline{\nabla}\partial_{t}\phi + \underline{w}n(\overline{\nabla}\phi\cdot\overline{\nabla}\phi - j\overline{\nabla}\phi_{\tau}\cdot\overline{\nabla}\phi_{\tau})) = 0$$
(5.6)

where j = 0 for instantaneous combustion and j = 1 for history-dependent combustion.

The most general solution of (5.6) (subject to hard-wall boundary conditions for the unsteady variables) can be written in the form of the following Fourier-Bessel series.

$$\phi = f_{00}(t) + \sum_{n=1}^{\infty} f_{0n}(t) J_0(S_{0n}r)$$

$$+\sum_{m=1}^{\infty}\sum_{n=1}^{\infty} (f_{mn}(t)\cos(m\theta)+g_{mn}(t)\sin(m\theta))J_{m}(S_{mn}r)$$
 (5.7)

Substituting (5.7) into (5.6) and applying the usual Galerkin orthogonalization procedure leads to an infinite

set of coupled ordinary differential equations governing the functions f_{mn} and g_{mn} . No solution can be obtained unless the series (5.7) is truncated so as to produce a finite set of equations. If the terms neglected are actually small, an accurate solution will result.

In this thesis attention will be focused on initial disturbances having the form of the first tangential mode. The simplest finite series contained in (5.7) capable of modeling the effect of quadradic nonlinearities in (5.6) is

$$\phi = f_0(t)J_0(S_{01}^r) + f_1(t)J_1(S_{11}^r)\cos\theta + f_2(t)J_2(S_{21}^r)\cos2\theta + g_1(t)J_1(S_{11}^r)\sin\theta + g_2(t)J_2(S_{21}^r)\sin2\theta.$$
 (5.8)

The Galerkin method then produces the following ordinary differential equations governing these five variables.

$$\ddot{f}_{0} + \underline{w}\dot{f}_{0} + S_{01}^{2}f_{0} + \varepsilon(C_{1}f_{0}\dot{f}_{0} + C_{2}(f_{1}\dot{f}_{1} + g_{1}\dot{g}_{1})$$

$$+ C_{3}(f_{2}\dot{f}_{2} + g_{2}\dot{g}_{2}) + \underline{w}n(C_{11}f_{0}^{2} + C_{12}(f_{1}^{2} + g_{1}^{2}) + C_{13}(f_{2}^{2} + g_{2}^{2})$$

$$-j(C_{11}f_{0\tau} + C_{12}(f_{1\tau}^{2} + g_{1\tau}^{2}) + C_{13}(f_{2\tau}^{2} + g_{2\tau}^{2})))) = 0$$

$$\ddot{f}_{1} + \underline{w}\dot{f}_{1} + S_{11}^{2}f_{1} + \varepsilon(C_{4}f_{0}\dot{f}_{1} + C_{5}f_{1}\dot{f}_{0} + C_{6}(g_{1}\dot{g}_{2} + f_{1}\dot{f}_{2})$$

$$+ C_{7}(g_{2}\dot{g}_{1} + f_{2}\dot{f}_{1}) + \underline{w}n(C_{14}f_{0}f_{1} + C_{15}(f_{1}f_{2} + g_{1}g_{2})$$

$$- j(C_{14}f_{0\tau}f_{1\tau} + C_{15}(f_{1\tau}f_{2\tau} + g_{1\tau}g_{2\tau})))) = 0$$

$$\ddot{f}_{2} + \dot{w}\dot{f}_{2} + \dot{S}_{21}^{2}f_{2} + \varepsilon(C_{8}f_{0}\dot{f}_{2} + C_{9}f_{2}\dot{f}_{0} + C_{10}(g_{1}\dot{g}_{1}-f_{1}\dot{f}_{1})$$

$$+ \underline{w}n(C_{16}(f_{1}^{2}-g_{1}^{2})+C_{17}f_{0}f_{2}-j(C_{16}(f_{1\tau}^{2}-g_{1\tau}^{2})+C_{17}f_{0\tau}f_{2\tau})))) = 0$$

$$\ddot{g}_{1} + \dot{w}\dot{g}_{1} + \dot{S}_{11}^{2}g_{1} + \varepsilon(C_{4}f_{0}\dot{g}_{1} + C_{5}g_{1}\dot{f}_{0} + C_{6}(f_{1}\dot{g}_{2}-g_{1}\dot{f}_{2})$$

$$+ C_{7}(g_{2}\dot{f}_{1}-f_{2}\dot{g}_{1}) + \underline{w}n(C_{14}f_{0}g_{1}+C_{15}(f_{1}g_{2}-g_{1}\dot{f}_{2})$$

$$- j(C_{14}f_{0\tau}g_{1\tau}+C_{15}(f_{1\tau}g_{2\tau}-g_{1\tau}f_{2\tau}))) = 0$$

$$\ddot{g}_{2} + \dot{w}\dot{g}_{2} + \dot{S}_{21}^{2}g_{2} + \varepsilon(C_{8}f_{0}\dot{g}_{2}+C_{9}g_{2}\dot{f}_{0}-C_{10}(g_{1}\dot{f}_{1})$$

$$+ f_{1}\dot{g}_{1}) + \underline{w}n(C_{17}f_{0}g_{2}+2C_{16}f_{1}g_{1}))) = 0$$

$$(5.9)$$

The coefficients C_i (i=1,2,...17) are integrals of Bessel functions which must be computed numerically. The details of this calculation are discussed in appendix A.

The stability boundaries in the (n,ϵ) plane were determined first. For a given value of \underline{w} , a value of ϵ was selected and solutions were obtained for various values of n. In each case, if the modal amplitudes exhibited growth with time, the value of n was decreased for the next run while if decay of modal amplitudes was observed, the

value of n was increased accordingly. A systematic iteration process was devised so that the computer could carry out these calculations without analyst intervention. In this way, one point on the stability boundary was Then a new value of ε was chosen and the established. iteration process was repeated to determine another point on the stability boundary. This procedure (which consumed a considerable amount of computer time) was continued until enough points had been found to establish the shape of the stability boundary. It should be pointed out that the amplitude of the 1T mode always initially decreases due to the fact that purely velocity-sensitive combustion instability is always linearly stable. Thus it is necessary to continue the calculation for a considerable period of time to determine whether a given set of conditions corresponds to nonlinear stability or instability.

Figures 2 and 3 show some typical stability boundaries for instantaneous combustion using the initial conditions

$$f_1(0) = 1$$
, $f_0(0) = f_2(0) = g_1(0) = g_2(0) = 0$
 $f_0(0) = f_1(0) = f_2(0) = g_1(0) = g_2(0) = 0$ (5.10)

and
$$f_1(0) = 1$$
, $f_0(0) = f_2(0) = g_1(0) = g_2(0) = 0$
 $g_1(0) = 1$, $f_0(0) = f_1(0) = f_2(0) = g_2(0) = 0$, (5.11)

respectively. The region below and to the left of the stability boundary is associated with stable conditions, while the region above and to the right is associated with unstable conditions.

In the case of linear acoustics, the first set of initial conditions would lead to a standing wave and the second set of initial conditions corresponds to a traveling wave. In both cases, it can be seen that the stability boundaries have roughly the forms of rectangular hyperbolas. As expected, increasing the steady-state burning rate reduces the value of n required to produce instability. This effect is somewhat more pronounced for the traveling waves than for the standing waves. There does not appear to be a distinct pattern to the results. Thus for $\underline{w} = 0.2$ standing waves are more stable than traveling waves while for $\underline{w} = 0.1$ traveling waves are more stable than standing waves.

Figures 4 and 5 show stability boundaries associated with initial conditions (5.10) and (5.11), respectively, for history-dependent combustion. It is apparent that the influence of the time delay parameter on the results for standing waves is much greater than on the results for traveling waves. Again, no clear pattern emerges from the results. Comparing Figures 2 and 4, and 3 and 5 shows that instantaneous combustion can be either more or less stable than history dependent combustion depending on the value of the time-delay parameter. Comparing Figures 4 and 5, it can be seen that standing waves can be either more or less

stable than standing waves for history dependent combustion. Figures 4 and 5 also illustrate the fact that increasing the amount of time delay can either increase or decrease the stability of the system. Thus, for both standing and traveling waves, $\tau = \pi$ corresponds to a more stable situation than either $\tau = 0.5\pi$ or $\tau = 1.5\pi$. The lack of patterns exhibited by these results emphasizes the need for numerical methods of the type developed during the present research. It appears that the only way to find out what will happen in a given situation is to solve the equations for that particular case. This matter will be discussed further subsequently.

The pressure perturbation can be calculated from the velocity potential perturbation by substituting equation (5.8) into (2.33c), expanding the result in a Fourier-Bessel series and retaining only the terms corresponding to the 1T, 2T and 1R modes to obtain

$$\begin{split} P &= - \gamma((d_{01}\dot{f}_0 + \varepsilon(d_{02}f_0^2 + d_{03}(f_1^2 + g_1^2) + d_{04}(f_2^2 + g_2^2)) \\ &+ d_{05}\dot{f}_0^2 + d_{06}(\dot{f}_1^2 + \dot{g}_1^2) + d_{07}(\dot{f}_2^2 + \dot{g}_2^2)))J_0(S_{01}r) \\ &+ ((d_{11}\dot{f}_1 + \varepsilon(d_{12}f_0f_1 + d_{13}(f_1f_2 + g_1g_2) + d_{14}\dot{f}_0\dot{f}_1) \\ &+ d_{15}(\dot{f}_1\dot{f}_2 + \dot{g}_1g_2)))\cos\theta \end{split}$$

$$+ (d_{11}\dot{g}_{1} + \epsilon(d_{12}f_{0}g_{1} + d_{13}(f_{1}g_{1} - g_{1}f_{2}) + d_{14}\dot{f}_{0}\dot{g}_{1}$$

$$+ d_{15}(\dot{f}_{1}\dot{g}_{2} - \dot{f}_{2}\dot{g}_{1})))\sin\theta)J_{1}(S_{11}r)$$

$$+ ((d_{21}\dot{f}_{2} + \epsilon(d_{22}f_{0}f_{2} + d_{23}(f_{1}^{2} - g_{1}^{2}) + d_{24}\dot{f}_{0}\dot{f}_{2}$$

$$+ d_{25}(\dot{f}_{1}^{2} - \dot{g}_{1}^{2}))\cos2\theta$$

$$+ (d_{21}\dot{g}_{1} + \epsilon(d_{22}f_{0}g_{2} + 2d_{23}f_{1}g_{1} + d_{24}\dot{f}_{0}\dot{g}_{2}$$

$$+ 2d_{25}\dot{f}_{1}g_{1})\sin2\theta)J_{2}(S_{21}r)).$$

$$(5.12)$$

The coefficients d_{0s} , d_{1s} , d_{2s} are calculated in the same fashion as those discussed previously and the values are listed in table 8 of the appendix.

Some typical results for wall pressure waveforms are presented in figures 6 through 37.

Figures 6 through 9 correspond to instantaneous combustion with ε = .05, \underline{w} = .1, n = 175, and initial conditions (5.10). This leads to a stable standing wave oscillation and the pressure can be observed to decrease gradually. However,by changing the interaction index to n = 220 (Figures 10-13) the pressure is seen to grow gradually. This is an unstable situation. In both cases the response is dominated by the 1T mode but distorted to some extent by the presence of the 2T and 1R components.

Figures 14 through 17 correspond to history-dependent

combustion with n = 345, while the other parameters are the same as before. This is a stable situation and the amplitude of the pressure slowly decays with time.

Figures 18 through 21 are obtained by changing the interaction index to n=352. This is an unstable situation as indicated by the growth of the pressure amplitude with time. In both of these cases the presence of the 2T and 1R components is much more noticable than it was in the first two situations.

Figures 22 through 25 correspond to instantaneous combustion with n = 180, ε = .05, \underline{w} = .1, and initial condition (5.11). This produces a traveling wave. The pressure is observed to be decreasing with time (a stable situation) while changing the interaction index n to 210 (Figures 26 through 29) causes the pressure to increase (an unstable situation). Figures 30 through 37 show the significance of the time delay function. Figures 30 through 33 are obtained by setting τ = π and n = 197. This is a stable case. The pressure is observed to decrease. Setting n = 199 (Figures 34 through 37), on the other hand, produces an unstable situation when the pressure increases very rapidly. In both of these situations the response appears to be dominated by 2T contribution to the pressure.

From these figures one can conclude that the pressure waveforms exhibit a strong second harmonic distortion and this distortion arises from the effect of the quadratic nonlinear terms. A variety of behaviors are possible

depending on the nature of the combustion process and the parametric values involved.

Chapter 6

ONE-DIMENSIONAL MODEL

In this chapter an annular combustion chamber with a gap width much smaller than the inner radius is considered. This geometry will henceforth be referred to as that of a narrow annulus. While this geometry is not of much practical interest, it is quite useful for the purposes of analysis. This is because only one space coordinate is needed to describe the problem. This makes a direct finite-difference numerical solution of the original partial differential equation feasible and also simplifies the algebra required to carry out the Galerkin modal analysis. In what follows three questions will be investigated. First, the effect of changing the numerical values of certain coefficients appearing in the governing equations for the modal amplitudes will be discussed. Second the Galerkin solution will be checked using a finite difference numerical solution of the complete equation. Third, numerical solutions of the complete wave equation using the vaporization-limited burning-rate law will be compared to similar solutions associated with the phenomenological burning-rate law employed in the previous chapter.

The appropriate wave equation for transverse wave motion in a narrow annulus can be obtained from (5.6) by

setting r=1 and $\vartheta_r=0$. To further simplify the results the parametric values $\gamma=1$ (isothermal process) and j=0 (instantenous burning response) will be employed. For this situation (5.6) simplifies to

$$\partial_{tt}\phi + \underline{w}\partial_{t}\phi - \partial_{\theta\theta}\phi + 2\varepsilon\partial_{\theta}\phi\partial_{\theta t}\phi + \underline{w}n\varepsilon(\partial_{\theta}\phi)^{2} = 0.$$
 (6.1)

To carry out the modal analysis, it is assumed that the potential function can be expressed as

$$\phi = f_1(t)\cos\theta + f_2(t)\cos2\theta + g_1(t)\sin\theta + g_2(t)\sin2\theta.$$
 (6.2)

Then applying the usual Galerkin orthogonalization procedure leads to

$$\ddot{f}_{1} + \Omega_{1}^{2}f_{1} + \underline{w}\dot{f}_{1} + \varepsilon A_{1}(f_{1}\dot{f}_{2} + f_{2}\dot{f}_{1} + g_{1}\dot{g}_{2} + g_{2}\dot{g}_{1})$$

$$+ \varepsilon \underline{w}nA_{2}(f_{1}f_{2} + g_{1}g_{2}) = 0$$

$$\ddot{f}_{2} + \Omega_{2}^{2}f_{2} + \underline{w}\dot{f}_{2} + \varepsilon A_{3}(g_{1}\dot{g}_{1} - f_{1}\dot{f}_{1}) + \varepsilon \underline{w}nA_{4}(g_{1}^{2} - f_{1}^{2}) = 0$$

$$\ddot{g}_{1} + \Omega_{1}^{2}g_{1} + \underline{w}\dot{g}_{1} + \varepsilon A_{1}(g_{2}\dot{f}_{1} + f_{1}\dot{g}_{2} - g_{1}\dot{f}_{2} - f_{2}\dot{g}_{1})$$

$$+ \varepsilon \underline{w}nA_{2}(f_{1}g_{2} - g_{1}f_{2}) = 0$$

$$\ddot{g}_{2} + \Omega_{2}^{2}g_{2} + \underline{w}g_{2} - \varepsilon A_{3}(g_{1}\dot{f}_{1} + f_{1}\dot{g}_{1}) - 2\varepsilon \underline{w}nA_{4}f_{1}g_{1} = 0$$
 (6.3)

where Ω_1 = 1, Ω_2 = 2, A_1 = 2, A_2 = 2, A_3 = 1 and A_4 = .5. The symbols Ω_1 , Ω_2 , A_1 , A_2 , A_3 and A_4 have been inserted to illustrate the effect of changing their numerical values.

McDonald (7) in his analysis of combustion—instability in an annulus found that for a given value of ε the value of n required to produce instability was approximately twice as high for standing waves as for traveling waves. The results discussed in the previous chapter for a full cylinder indicated no such relationship. Equations (6.3) were employed in an attempt to determine the factors which have a significant effect on the stability boundary. Several calculations were made and a representative sample of the data thus obtained is presented in Tables 1 and 2. It was determined by McDonald that the terms representing gas-dynamic nonlinearities, had a small qualitative effect on stability calculations. Thus the coefficients A_1 and A_3 were held fixed during these calculations.

The entries in the first two lines of each table were computed using the correct equation for an annulus. It can be seen that the standing wave is twice as stable as the traveling wave, in agreement with the results of McDonald. The third and fourth lines in each table were computed by changing A_{μ} from .5 to .77 (This makes the ratio A_{μ}/A_{3} the same as the ratio of the corresponding terms in the governing equations for the full cylinder.). It can be seen that this lowers the stability limit in all cases but does not alter the fact that an initial disturbance in the

Table 1
Stability Limits for Standing Wave in Annular Combustor

n	^Ω 2	A ₄	ε
91 46 74 37 735 379 595 307	2 2 2 1.66 1.66 1.66	•5 •77 •77 •5 •77	.05 .1 .05 .1 .05 .1

Table 2
Stability Limits for Traveling Wave in Annular Combustor

n	${\mathfrak a}^5$	Ац	ε
44 22 36 18 66 31 54 26	2 2 2 1.66 1.66 1.66	.5 .77 .77 .5 .5 .77	.05 .1 .05 .1 .05 .1

form of a standing wave is twice as stable as one in the form of a traveling wave.

The fifth and sixth lines in Tables 1 and 2 are found by restoring $\rm A_4$ to its original value .5 and changing $\rm \Omega_2$ from 2.00 to 1.66 (This makes the ratio $\rm \Omega_2/\Omega_1$ equal to the ratio $\rm S_{21}/\rm S_{11}$ associated with the full cylinder.). For this situation it can be seen that the standing wave is approximately ten times as stable as the traveling wave. Furthermore, the effect of the value of $\rm \Omega_2$ on the location of stability boundary is much greater for a standing-wave motion than for a traveling-wave motion. The last two lines are associated with implementing both of the changes discussed above simultaneously. These results confirm that the change in $\rm A_4$ lowers the stability limit in all cases but does not affect the relative stability of standing-wave and traveling-wave disturbances.

The data presented above indicate that standing waves will be twice as stable as traveling waves only under very special circumstances ($\Omega_2/\Omega_1 \simeq 2$). There is no reason to expect this to be a characteristic of other systems exhibiting combustion instability as, in fact, it is not for a full cylinder.

In this section, a finite different procedure is employed to generate a set of second order differential equations. The central difference formulas

$$\partial_{\theta} \phi_{i} = (\phi_{i+1} - \phi_{i-1})/(2\Delta\theta)$$

$$\partial_{\theta\theta} \phi_{i} = (\phi_{i+1} - 2\phi_{i} + \phi_{i-1})/(\Delta\theta)^{2}$$
 (6.4)

are used to produce the following set of governing equations

$$\ddot{\phi}_{i} = -\underline{w}\dot{\phi}_{i} + (\phi_{i+1} - 2\phi_{i} + \phi_{i-1})(1 - \varepsilon(\gamma - 1)\dot{\phi}_{i})/(\Delta\theta)^{2}$$

$$- \varepsilon(\phi_{i+1} - \phi_{i-1})(\dot{\phi}_{i+1} - \dot{\phi}_{i-1})/(2\Delta\theta)^{2}$$

$$- \varepsilon w_{i} \qquad 2 < i < N - 1 \qquad (6.5)$$

where N is the number of points employed. A forward difference formula

$$\theta_{\theta}^{\phi_1} = (-3\phi_1 + 4\phi_2 - \phi_3)/(2\Delta\theta)$$
 (6.6)

and backward difference formula

$$\theta_{\theta} \phi_{N} = (\phi_{N-2} - 4\phi_{N-1} + 3\phi_{N})/(2\Delta\theta)$$
 (6.7)

are used for the boundary conditions

$$a_{\theta} \phi(0) = 0, \qquad a_{\theta} \phi(\pi) = 0.$$
(6.8)

Hence, along the boundaries, (6.5) becomes

$$\dot{\phi}_2 = -\underline{w}\dot{\phi}_2 + 2(\dot{\phi}_3 - \dot{\phi}_2)(1 - \varepsilon(\gamma - 1)\dot{\phi}_2)/(3\Delta\theta^2)$$

$$-8\varepsilon(\phi_3-\phi_2)(\phi_3-\phi_2)/(9\Delta\theta^2)-\varepsilon w_2$$

and
$$\phi_{N-1} = -\underline{w}\phi_{N-1} + 2(\phi_{N-2} - \phi_{N-1})(1-\epsilon(\gamma-1)\phi_{N-1})/(3\Delta\theta^2)$$

$$-8\varepsilon(\phi_{N-1} - \phi_{N-2})(\phi_{N-1} - \phi_{N-2})/(9\Delta\theta^{2}) - \varepsilon w_{N-1}. \quad (6.9)$$

For the phenomenological model, the burning rate function becomes

$$w_i = n\underline{w}(\phi_{i+1} - \phi_{i-1})^2/(4\Delta\theta^2)$$
 2 < i < N-1. (6.10)

Combining (6.8) and (6.10) one obtains

$$w_2 = 4n\underline{w}(\phi_3 - \phi_2)^2/(9\Delta\theta^2)$$

$$w_{N-1} = 4n\underline{w}(\phi_{N-1} - \phi_{N-2})^2/(9\Delta\theta^2)$$
 (6.11)

respectively.

For the vaporization model, the burning rate function becomes

$$w_{i} = \underline{w}((1 - \epsilon \phi_{i}/2)(1 + (\phi_{i+1})^{2})^{1/4} - 1)/\epsilon^{2} \qquad 2 < i < N-1$$
 (6.12)

and, along the boundary, yields

$$w_2 = \underline{w}((1 - \frac{1}{2}\epsilon\dot{\phi}_2)(1 + (2(\phi_3 - \phi_2)/(3\Delta\theta u_L))^2)^{\frac{1}{4}} - 1)/\epsilon^2$$

$$w_{N-1} = \underline{w} ((1 - \frac{1}{2} \epsilon \phi_{N-1}) (1 + (2(\phi_{N-1} - \phi_{N-2}) / (3\Delta \theta u_L))^2)^{\frac{1}{4}} - 1) / \epsilon^2, \quad (6.13)$$

accordingly.

Using either the phenomonological or the vaporization-limited combustion model,(6.5) and (6.9) can be solved by the Runge-Kutta method to obtain ϕ_i , $i=2,3,\ldots$, N-1. Then the modal amplitudes can be obtained by using the Fourier-cosine transform

$$f_{i}(t) = 2 \int_{0}^{\pi} \phi \cos i\theta d\theta / \pi. \qquad (6.14)$$

These integrals must be computed numerically since ϕ is known only at the grid points.

Some typical results for standing waves are presented in Table 3. They are intended to illustrate the accuracy of the two-term Galerkin solution and to compare the results associated with the phenomenological and vaporization-limited combustion laws. The column labeled PG indicates the results with a two-term Galerkin solution using the phenomenological model, the column labeled PF contains the results of a finite-difference solution of these equations, and the column labeled VF presents results of a finite difference solution using the vaporization-limited model.

Table 3

Stability	Limits	For	Phenomenological	and
Vaporiza	ation-Li	Lmite	d Combustion Mode	1

	PG	PF	VF
ε	n	n	u _L
.1 .05 .01	46 91 452	28 56 286	1.3 2.1 5.2

It can be seen that the order of magnitude of the interaction index n required for instability for both the finite difference and Galerkin methods are roughly the same but the stability boundaries predicted by the Galerkin method are approximately twice as high as those predicted by the Finite-difference method. It is to be expected that the Galerkin method will lead to higher stability limits than the use of an exact solution procedure. This can be explained as follows. The instability mechanism is basically a feedback process. Consider the form of (6.3) associated with standing waves in an annulus. This is

$$\ddot{f}_{1} + \underline{w}\dot{f}_{1} + f_{1} + 2\varepsilon(f_{1}\dot{f}_{2} + f_{2}\dot{f}_{1}) + 2\varepsilon\underline{w}nf_{1}f_{2} = 0$$

$$\ddot{f}_{2} + \underline{w}\dot{f}_{2} + 4f_{2} - \varepsilon f_{1}\dot{f}_{1} - \frac{1}{2}\varepsilon\underline{w}nf_{1}^{2} = 0. \tag{6.14}$$

For clarity the gas-dynamic nonlinearities will be neglected to give

$$\ddot{f}_{1} + \underline{w}\dot{f}_{1} + f_{1} + 2\varepsilon \underline{w}nf_{1}f_{2} = 0$$

$$\ddot{f}_{2} + \underline{w}\dot{f}_{2} + 4f_{2} - \frac{1}{2}\varepsilon \underline{w}nf_{1}^{2} = 0. \tag{6.15}$$

A one-term Galerkin solution would lead to the equation

$$\ddot{f}_{1} + \underline{w}\dot{f}_{1} + f_{1} = 0.$$

This would indicate unconditional stability. Instability can arise only when the presence of the last term in (6.15b) causes f_2 to grow and this then causes f_1 to grow because of the last term in (6.15a). The growth of f, also produces the growth of higher modes which in turn causes additional growth of f_1 . The contributions of these higher modes are neglected in a two-term Galerkin analysis and thus the energy input due to unsteady burning is underestimated. For this reason the Galerkin analysis will overestimate the stability of the system. This overestimation will decrease as the number of terms retained is increased. An example of this can be seen by comparing the one- and two-term analyses. A one-term solution predicts that the system is always stable. A two-term solution predicts the correct qualitative behavior but overestimates the system's stability by

roughly a factor of two. It is to be expected that further increases in accuracy can be obtained by keeping more terms in the assumed solution but that this will not seriously affect the quatitative prediction.

In Chapter 5, it was found that equating $dw/du_{\rm L}^2$ the phenomenological and vaporization-limited combustion laws at u^2 = 0 lead to the formula

$$n = 1/(4 \varepsilon^2 u_L^2)$$
. (6.16)

Assuming that the actual data can be fitted to a formula of the form

$$n = C/(4 \varepsilon^2 u_L^2)$$
 (6.17)

the data give in Table 3 provide an opportunity to estimate C. The results are shown below.

ε	.1	.05	.01	
С	1.89	2.47	3.09	

It appears that a value of C = 2.5 would give acceptable accuracy over this range of ϵ .

For a given value of u_L the n computed from (6.16) will overestimate the burning rate. Thus it might be thought that C should be less than unity. The stability criterion was, however, based on the magnitude of the modal amplitudes. Inspection of Tables 4 and 5 shows that the

vaporization-limited combustion model involves the higher modes to a greater extent than does the phenomenological combustion model. Thus, a given value of \mathbf{u}_L can be associated with lower individual modal amplitudes in the former case than in the latter. These two effects interact and calculation shows that C is greater than unity in this range of ϵ .

MODAL AMPLITUDE FOR PHENOMENLOGICAL MODEL IN ANNULAR COMBUSTOR

			. ~ ~ ~ ~ ~ .						
7	MODE 1	MODE 2	MODE 3	MODE 4	MODE 5	MODE 6	MODE 7	MODE 8	
05050505050505050505050505050505050505	077709175375300335547410761297172745682290278541324 036727255155498687884732292971727456883129299027854124 00000000000000000000000000000000000	0.000000000000000000000000000000000000	0208960584902330435859724361644669906982553231 00000001222001330224456907740698125955231 000000000000000000000000000000000000	00001132490110705704856050898977042682395183034744667 0000000011070570048560500552744367 000000000011070570048560552744367 000000000000000000000000000000000000	0000001123335670033458889013333444433320997417333166461510 000000000001010118889013333444433320997417333166466760 0000000000001010118889013333444433320997417333166466760 0000000000000000000000000000000	00000000121315154488115025737220327963265865828517 000000000000000000110010111202020202020	-0.000000122211446741600112834427074922093331 -0.0000000122211446741600112834427074922093331 -0.00000000000000000000000000000000000	00000000001101222203552266415851587160809366528222 000000000000000000000000000000000	

TABLE 5

MODAL AMPLITUDE FOR VAPORIZATION MODEL
IN ANNULAR COMBUSTOR

Chapter 7

CONCLUSIONS

The primary objective of this study was the development of a new analytical technique to be used in the solution of nonlinear velocity-sensitive combustion instability problems. Such a method should be relatively easy to apply and should require relatively little computation time.

In an attempt to achieve this aim, the orthogonal collocation method was investigated first. However, it was found that the results were heavily dependent on the location of the collocation points and characteristics of the equations. Therefore, the method was rejected as unreliable.

Next, the Galerkin method, which has proved to be very successful in analysis of the pressure sensitive combustion instability, was considered. This method proved to work very well. It was found that the pressure waveforms exhibit a strong second harmonic distortion and a variety of behaviors are possible depending on the nature of the combustion process and the parametric values involved.

Finally, a one-dimensional model provided further insight into the problem by allowing a comparison of Galerkin solutions with more exact finite-difference computations.

Some major conclusions of this research are as follows. (1) The form of (2.33a) is somewhat insensitive to the specific assumptions used to derive it. This is demonstrated by the fact that Powell (1) developed an equation of a very similar form using a different set of assumptions. (2) The orthogonal collocation method is unsuited to solution of problems of the type under discussion here. (3) The Galerkin method is well suited to the solution of such problems. (4) Stability boundaries and pressure wave forms appear to be highly dependent on the parameters of the problem (interaction index, time delay, steady-state burning rate, etc.). Furthermore, there appears to be no clear pattern to the computed results. (5) The phenomenological burning-rate law employed in the majority of the work discussed predicts results which are qualitatively similar to those associated with the vaporization-limited burning-rate law used by previous investigators. The phenomenological law, furthermore, can be used in conjunction with the Galerkin method while the vaporization-limited law cannot. (6) The computer programs developed in the course of this work can be employed to determine stability boundaries and pressure waveforms without the expenditure of excessive computer This is important because the lack of clear trends discussed above make an individual analysis of each situation desirable.

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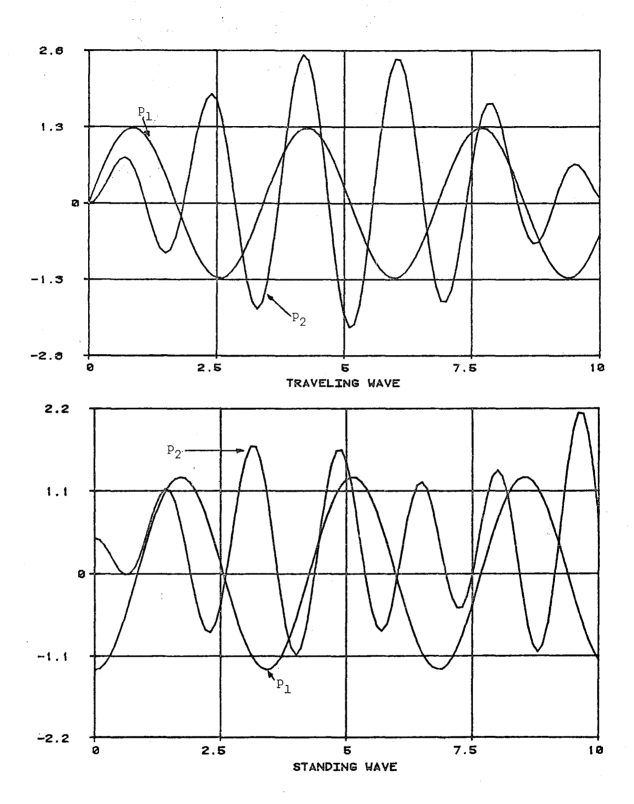


Figure 1. Pressure functions for standing and traveling waves

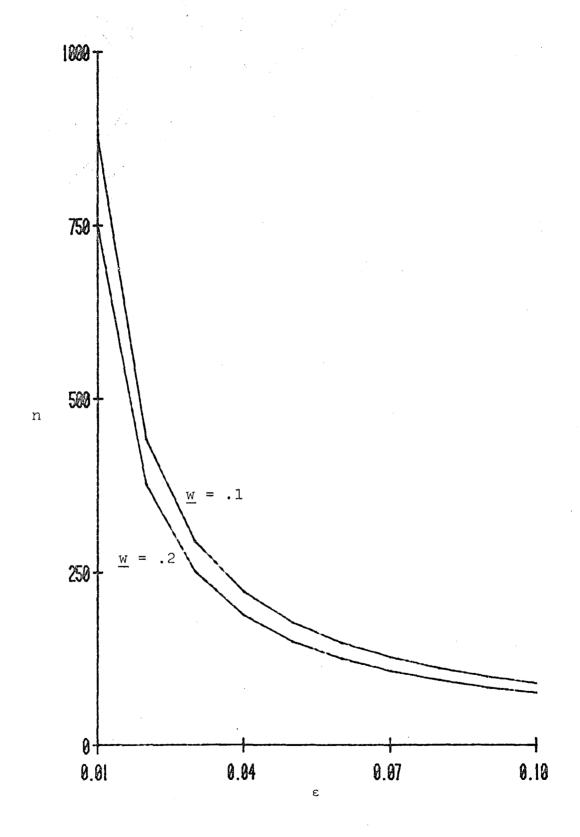


Figure 2. Stability Boundary for Standing Waves

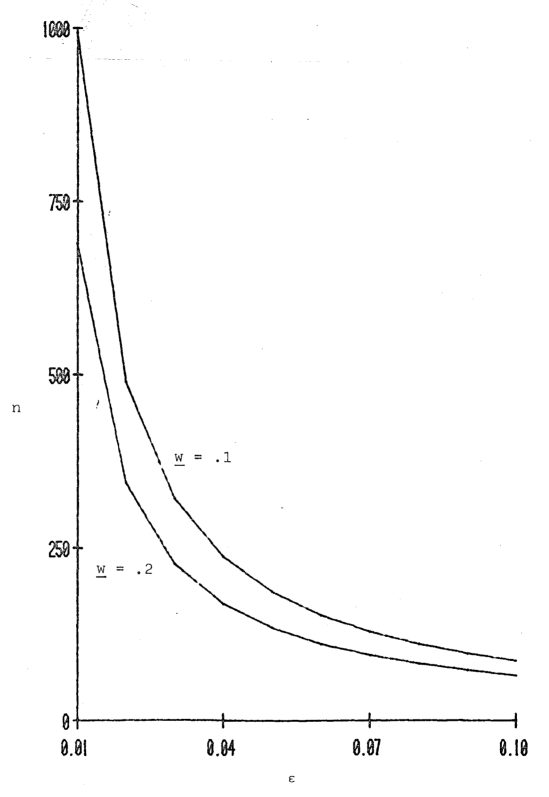


Figure 3. Stability Boundary for Traveling Waves

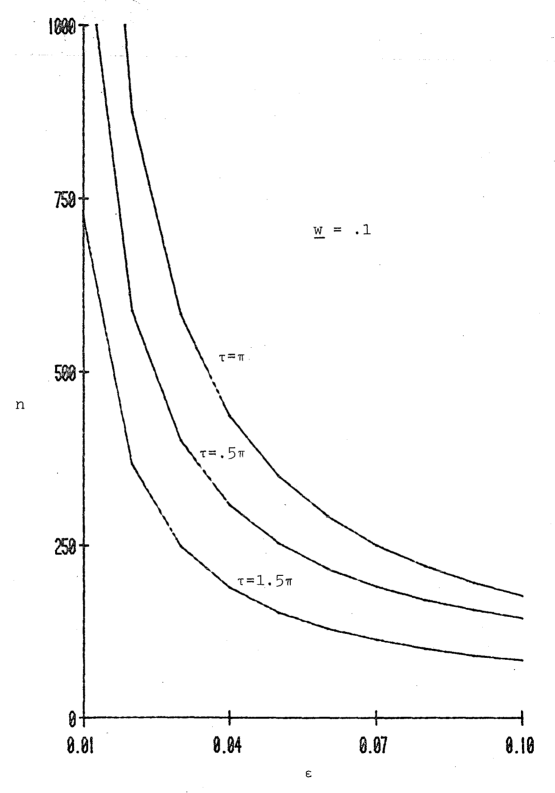


Figure 4. Stability Boundary for Standing Waves

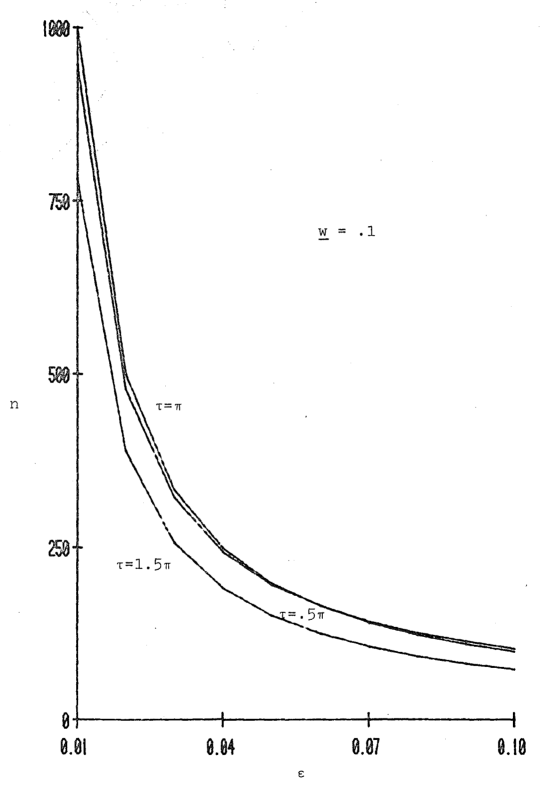


Figure 5. Stability Boundary for Traveling Waves

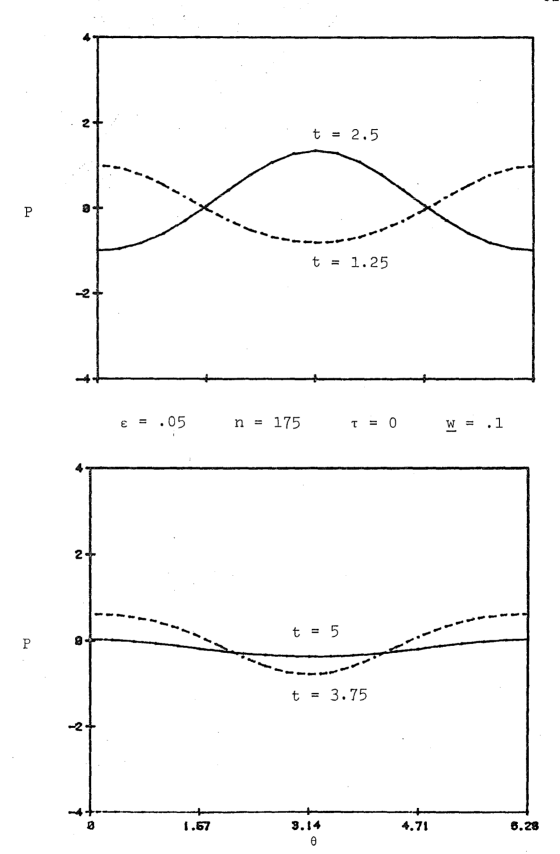


Figure 6. The Wall Pressure Waveforms for Standing Waves

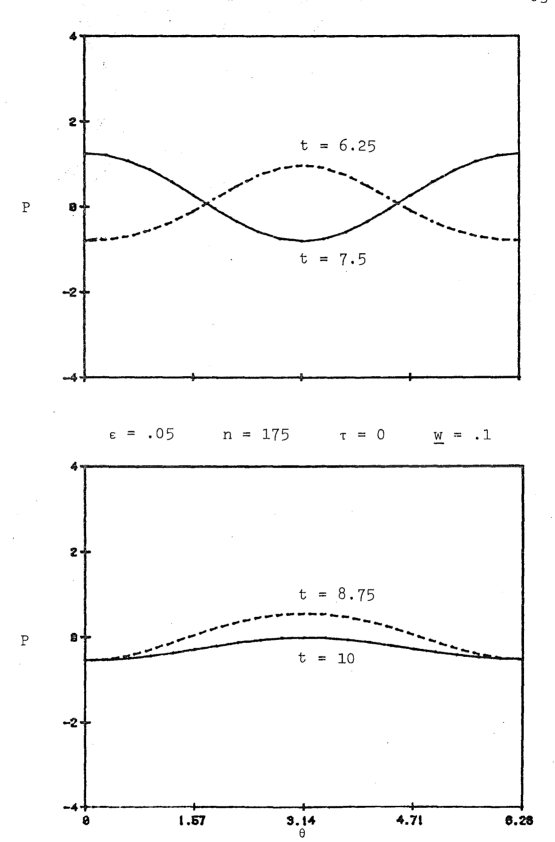


Figure 7. The Wall Pressure Waveforms for Standing Waves

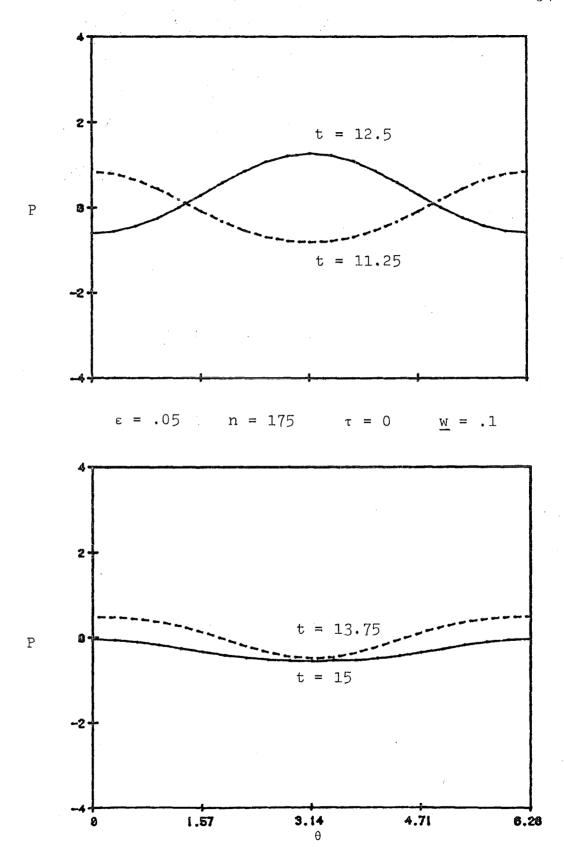


Figure 8. The Wall Pressure Waveforms for Standing Waves

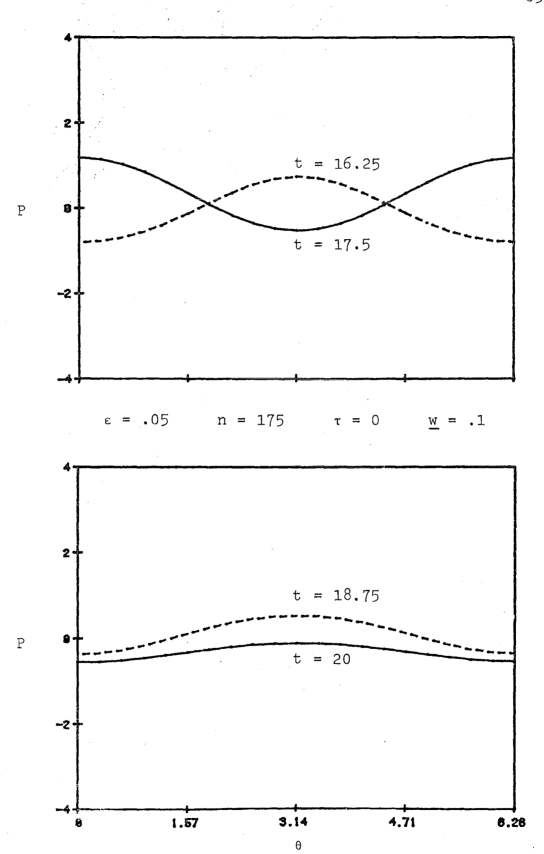


Figure 9. The Wall Pressure Waveforms for Standing Waves

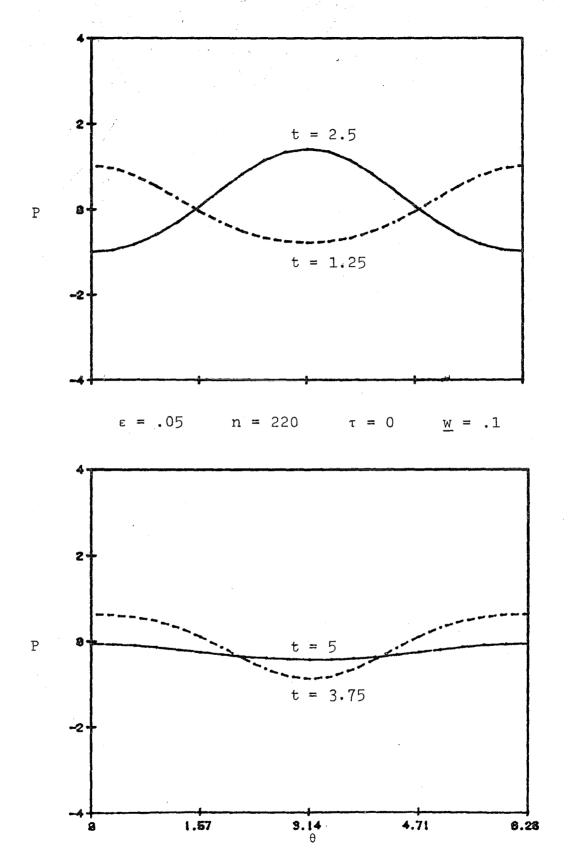


Figure 10. The Wall Pressure Waveforms for Standing Waves

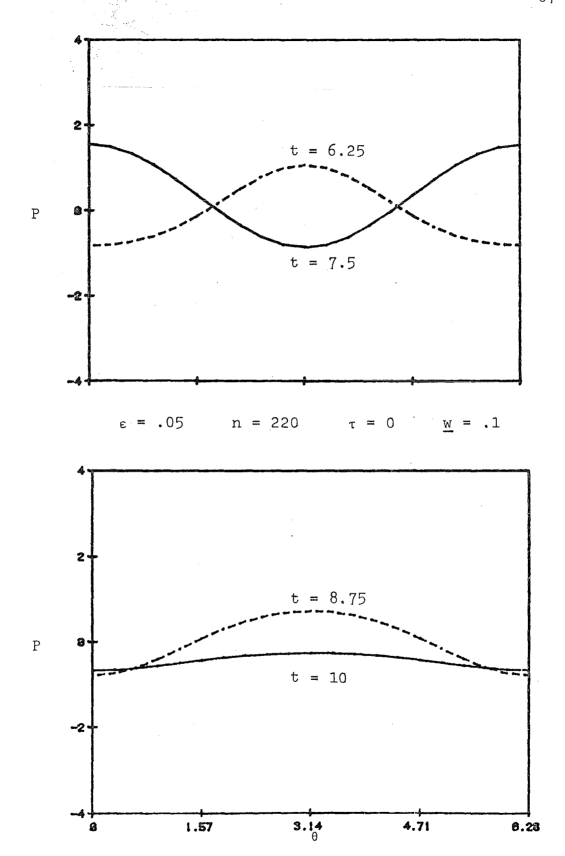


Figure 11. The Wall Pressure Waveforms for Standing Waves

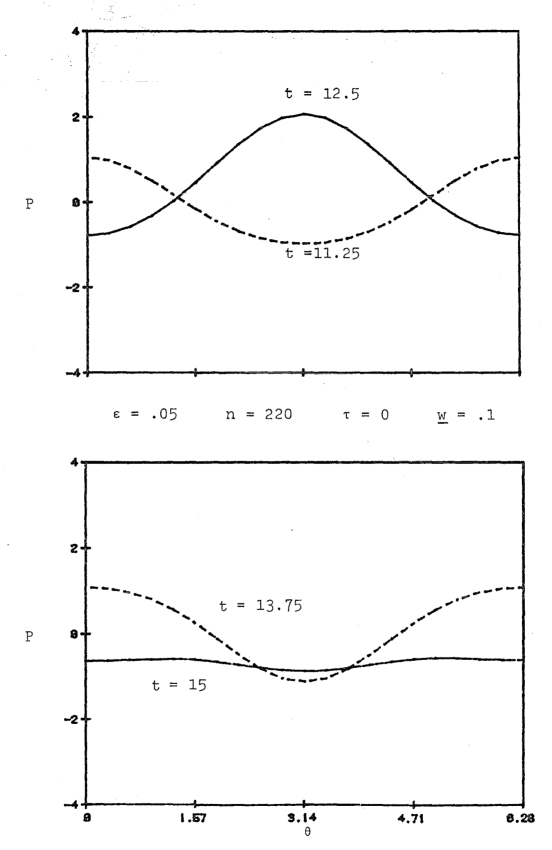


Figure 12. The Wall Pressure Waveforms for Standing Waves

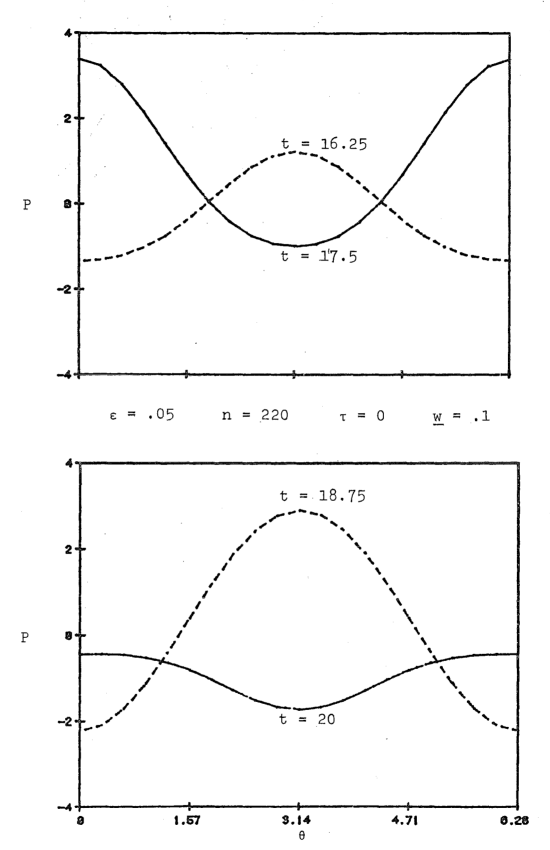


Figure 13. The Wall Pressure Waveforms for Standing Waves

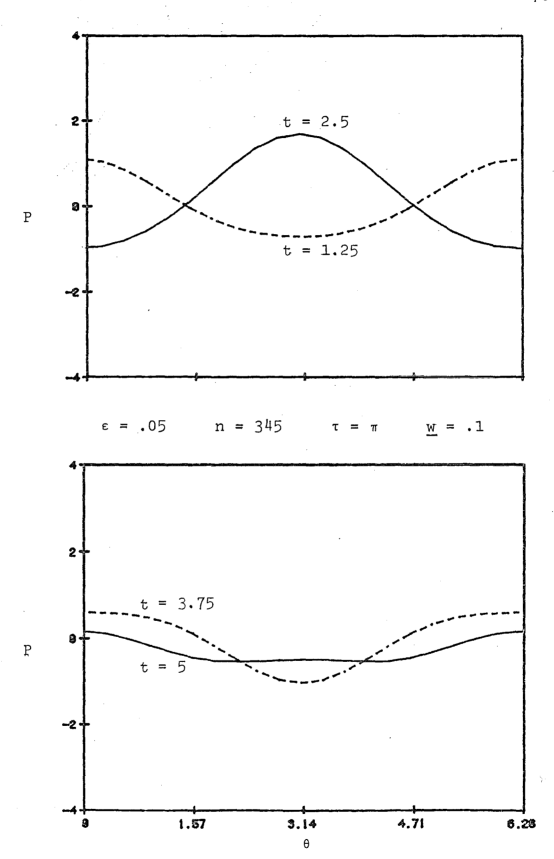


Figure 14. The Wall Pressure Waveforms for Standing Waves

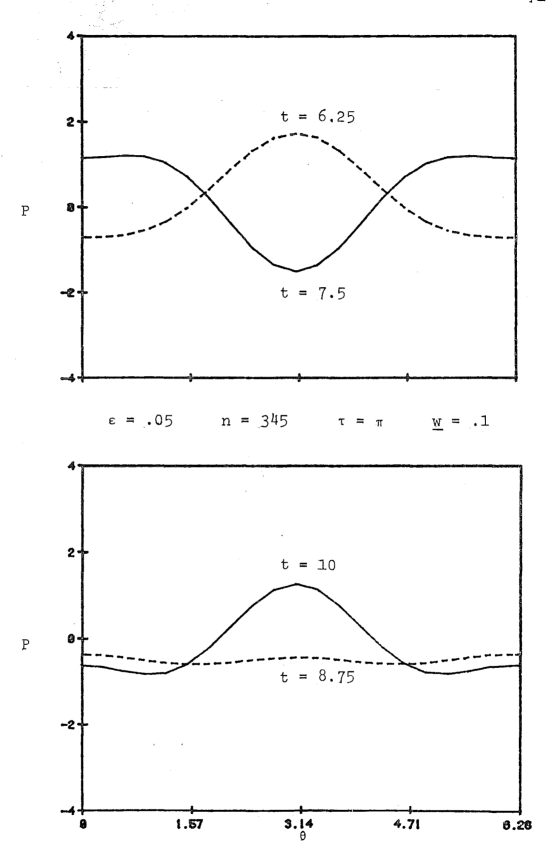


Figure 15. The Wall Pressure Waveforms for Standing Waves

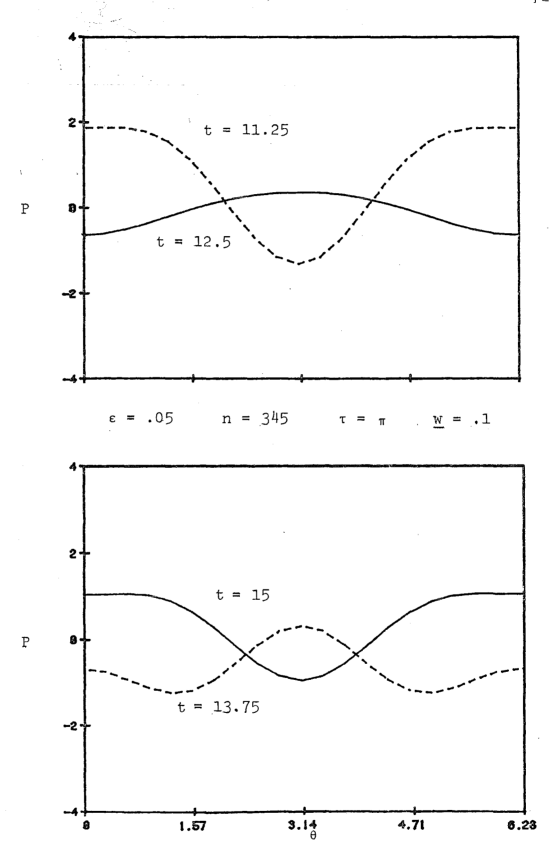


Figure 16. The Wall Pressure Waveforms for Standing Waves

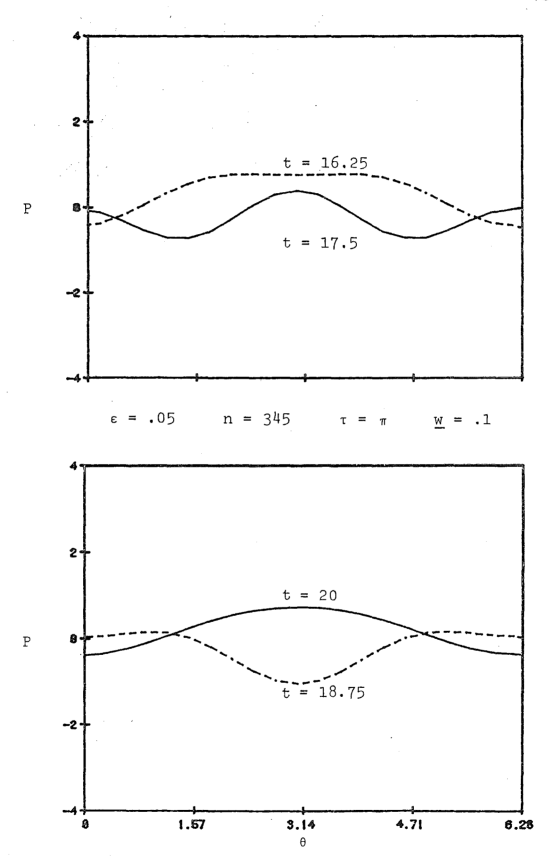


Figure 17. The Wall Pressure Waveforms for Standing Waves

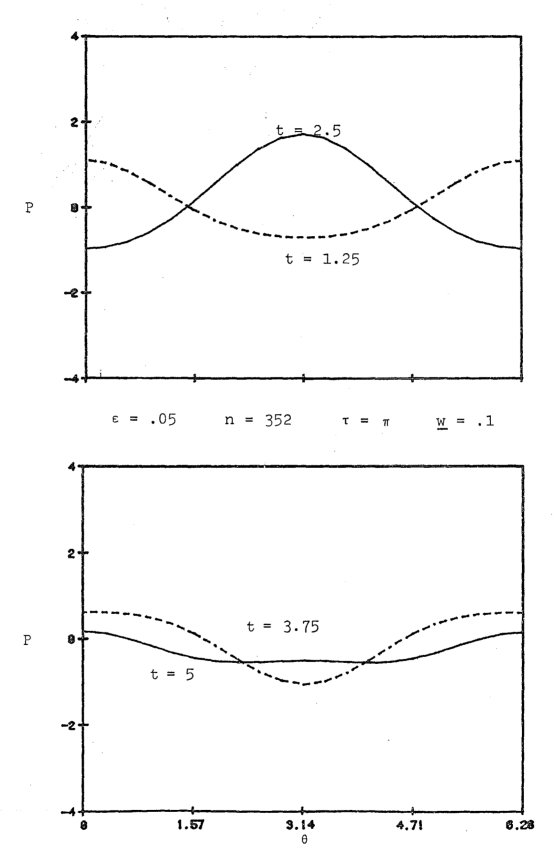


Figure 18. The Wall Pressure Waveforms for Standing Waves

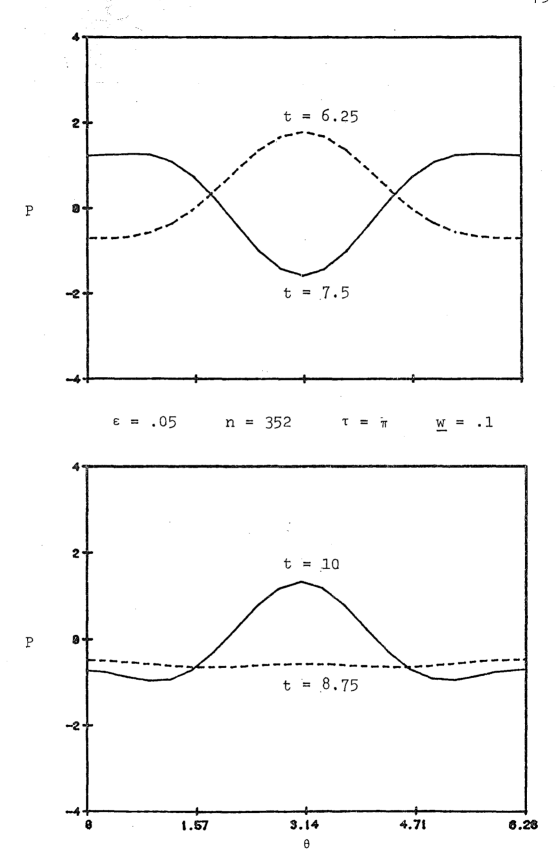


Figure 19. The Wall Pressure Waveforms for Standing Waves

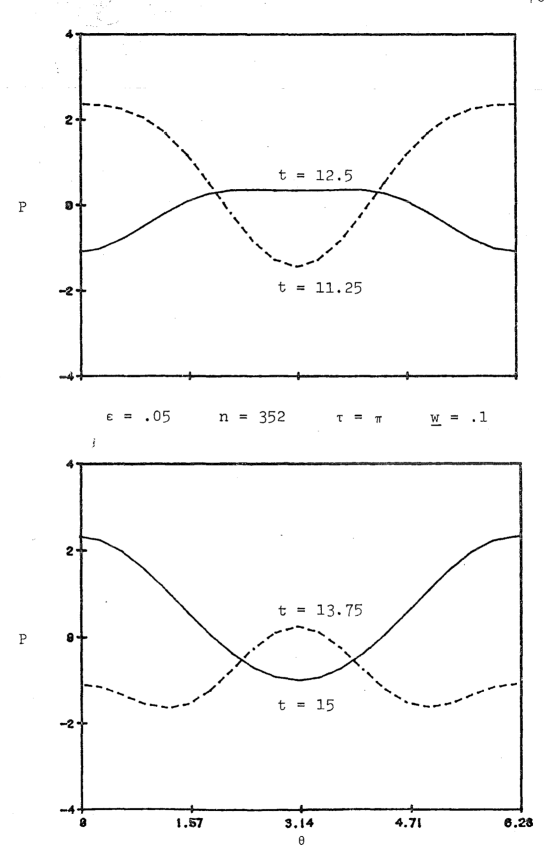


Figure 20. The Wall Pressure Waveforms for Standing Waves

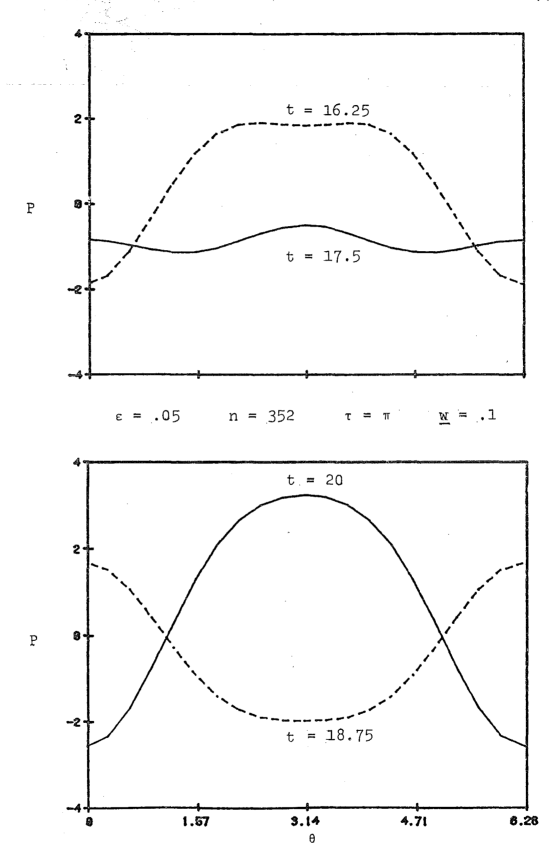


Figure 21. The Wall Pressure Waveforms for Standing Waves

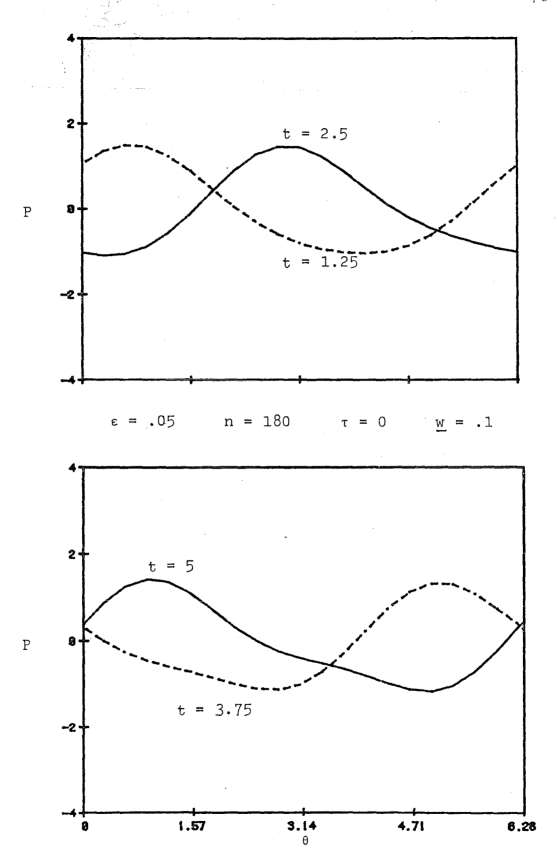


Figure 22. The Wall Pressure Waveforms for Traveling Waves

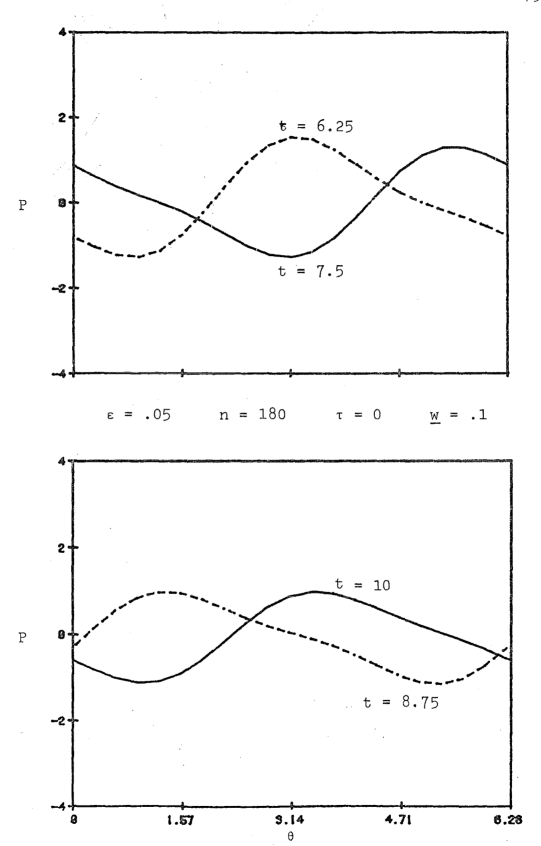


Figure 23. The Wall Pressure Waveforms for Traveling Waves

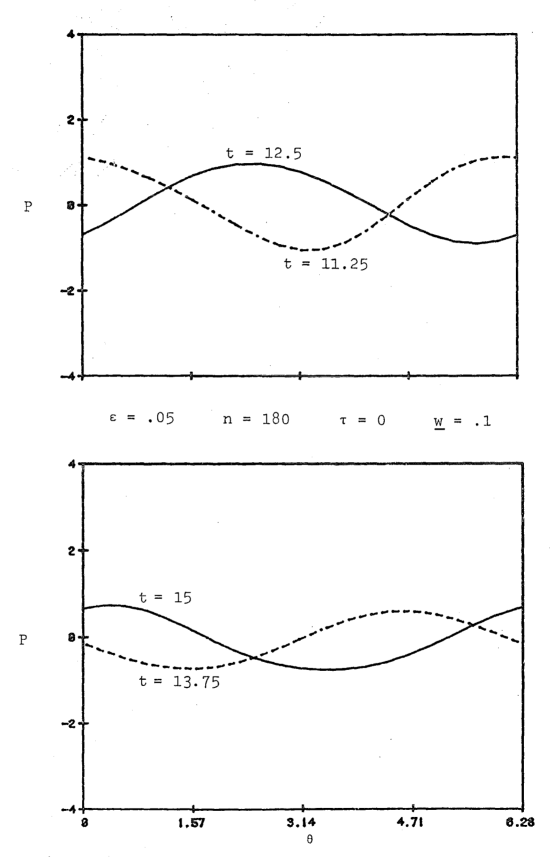


Figure 24. The Wall Pressure Waveforms for Traveling Waves

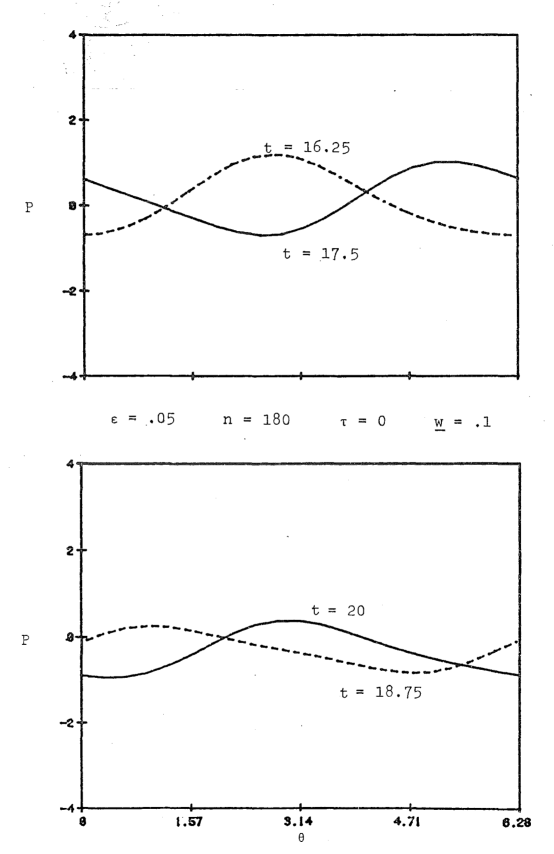


Figure 25. The Wall Pressure Waveforms for Traveling Waves

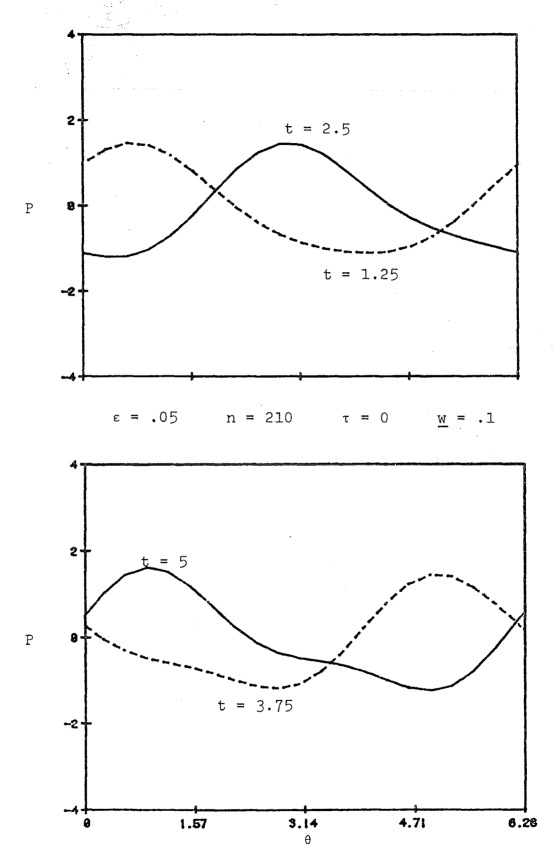


Figure 26. The Wall pressure Waveforms for Traveling Waves

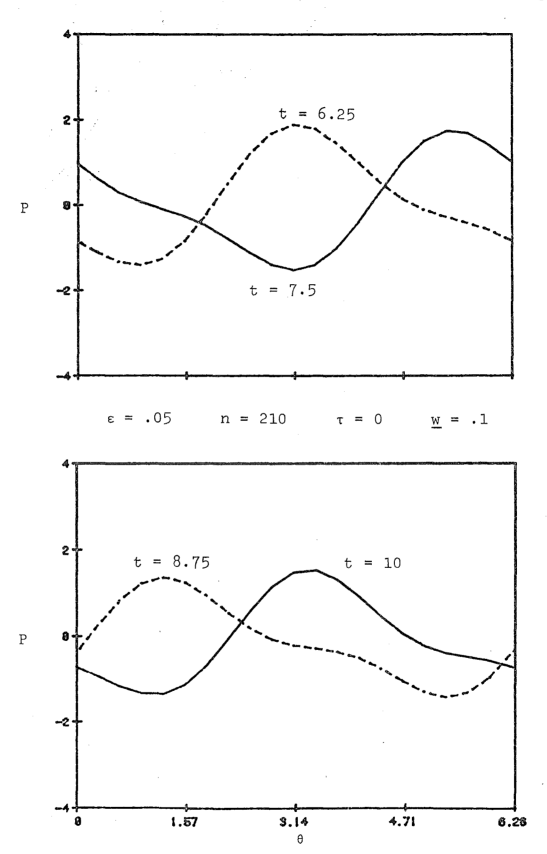


Figure 27. The Wall Pressure Waveforms for Traveling Waves

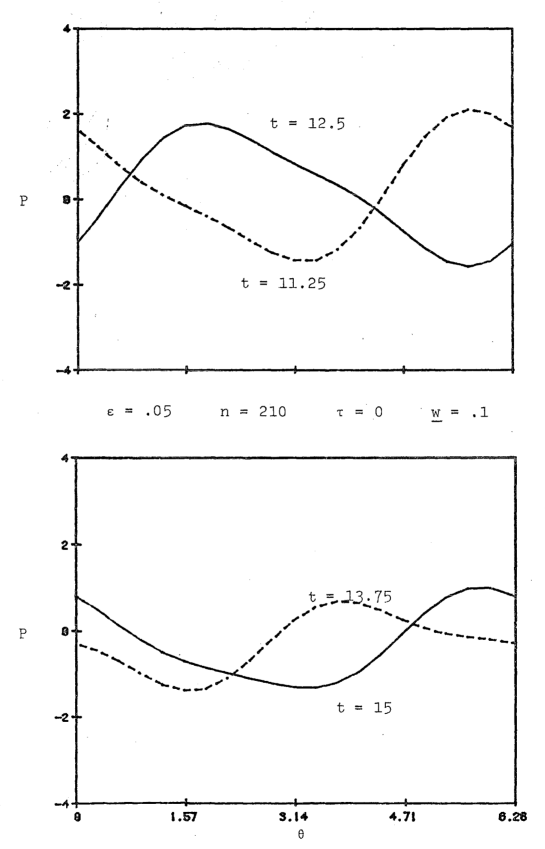


Figure 28. The Wall Pressure Waveforms for Traveling Waves

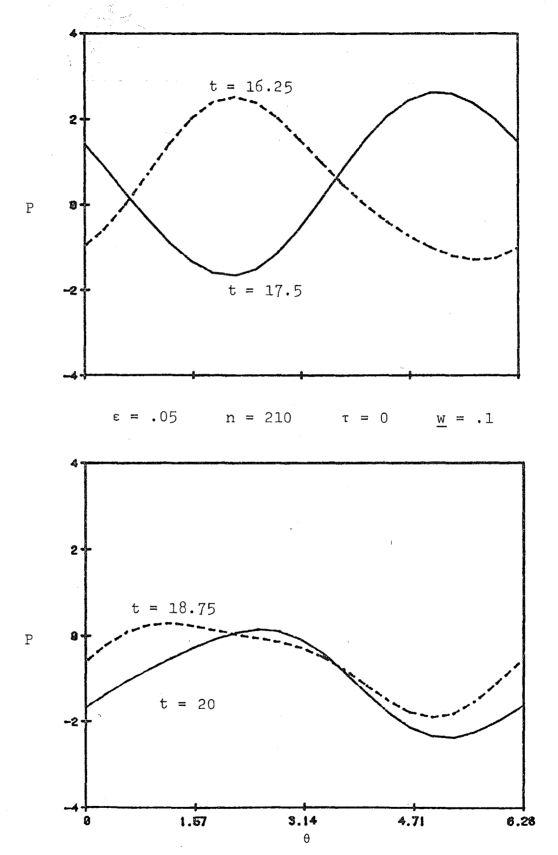


Figure 29. The Wall Pressure Waveforms for Traveling Waves

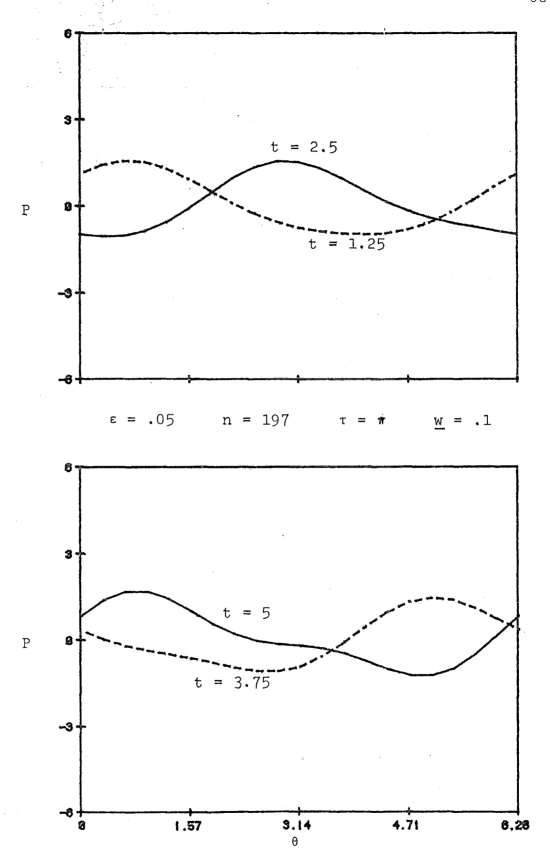


Figure 30. The Wall Pressure Waveforms for Traveling Waves

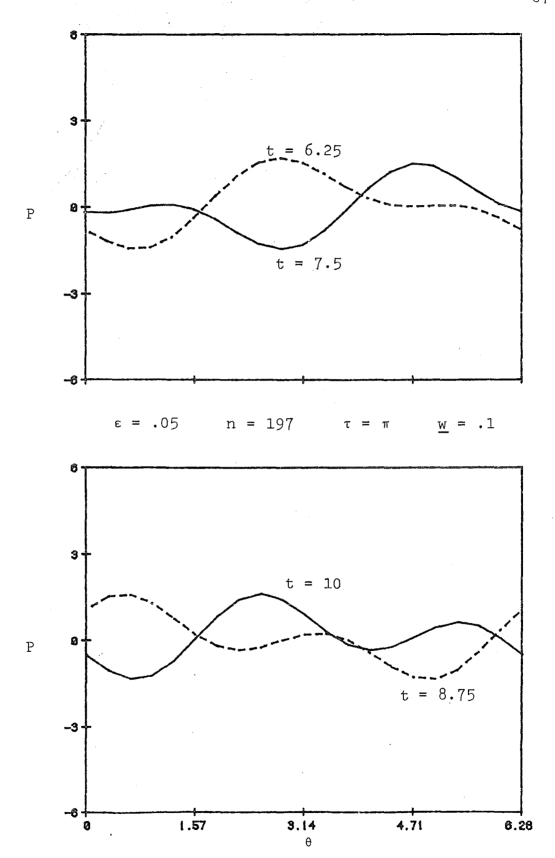


Figure 31. The Wall Pressure Waveforms for Traveling Waves

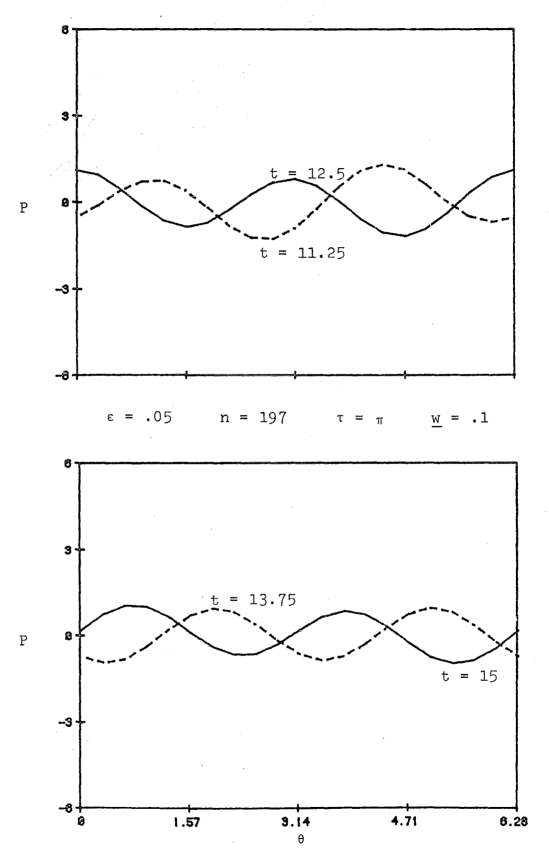


Figure 32. The Wall Pressure Waveforms for Traveling Waves

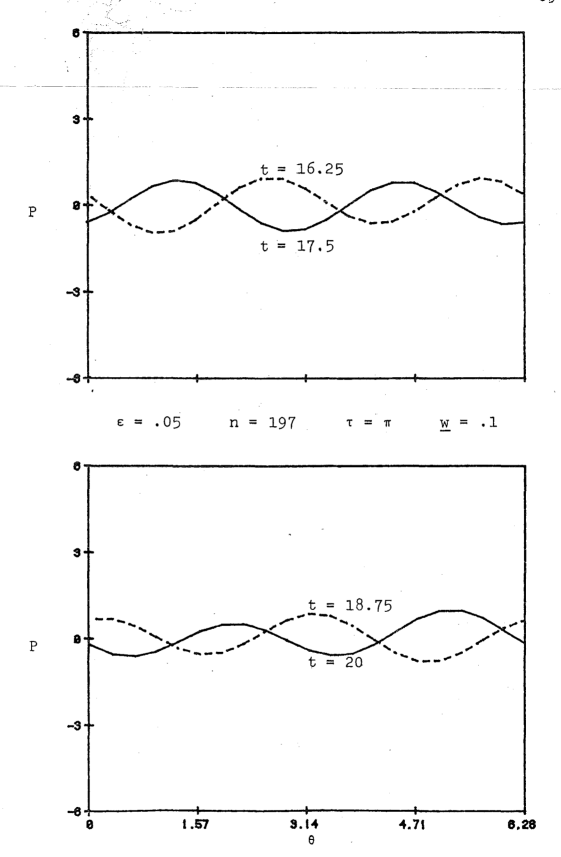


Figure 33. The Wall Pressure Waveforms for Traveling Waves

. . . .

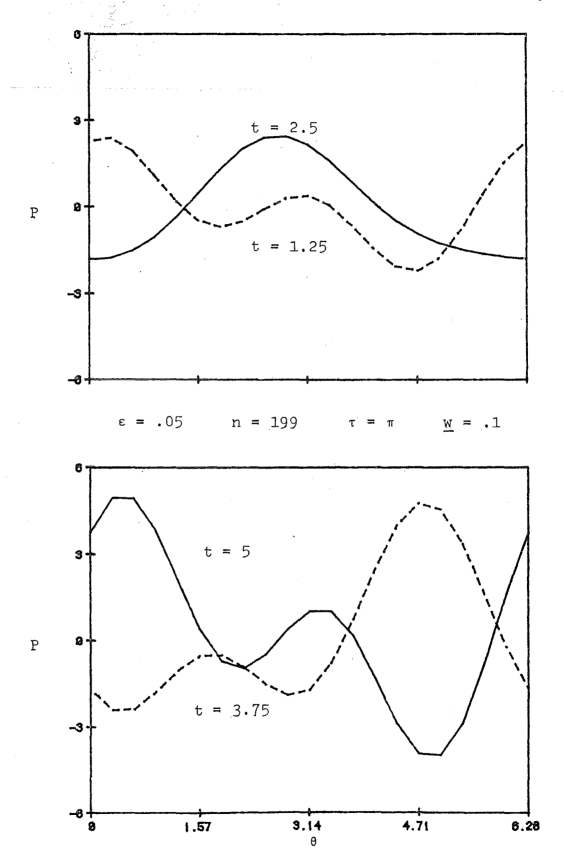


Figure 34. The Wall Pressure Waveforms for Traveling Waves

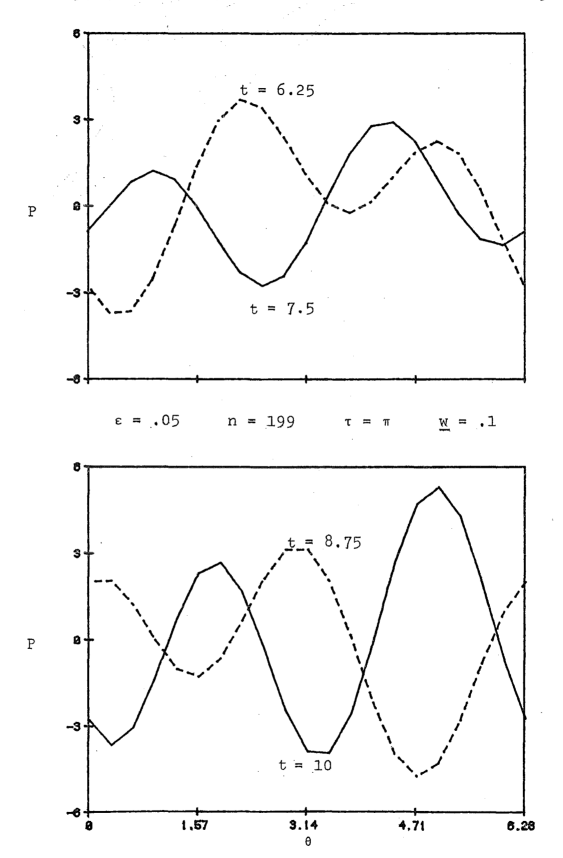


Figure 35. The Wall Pressure Waveforms for Traveling Waves

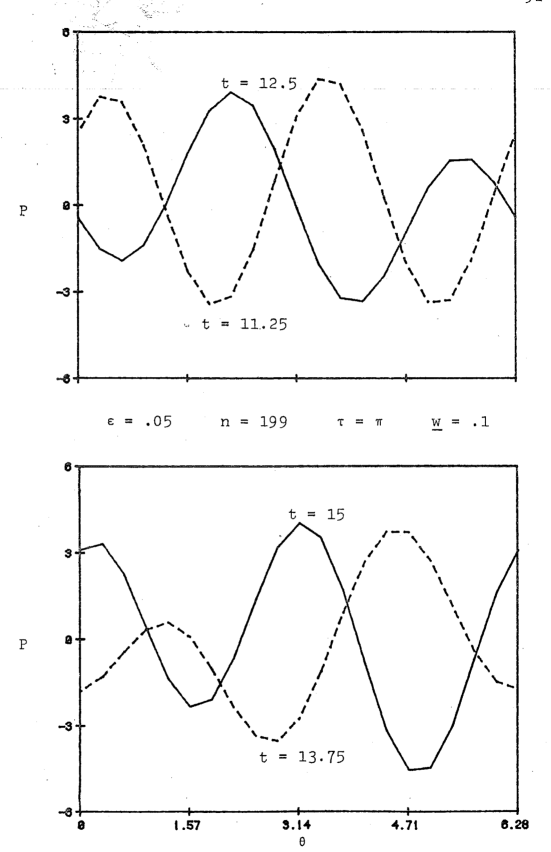


Figure 36. The Wall Pressure Waveforms for Traveling Waves

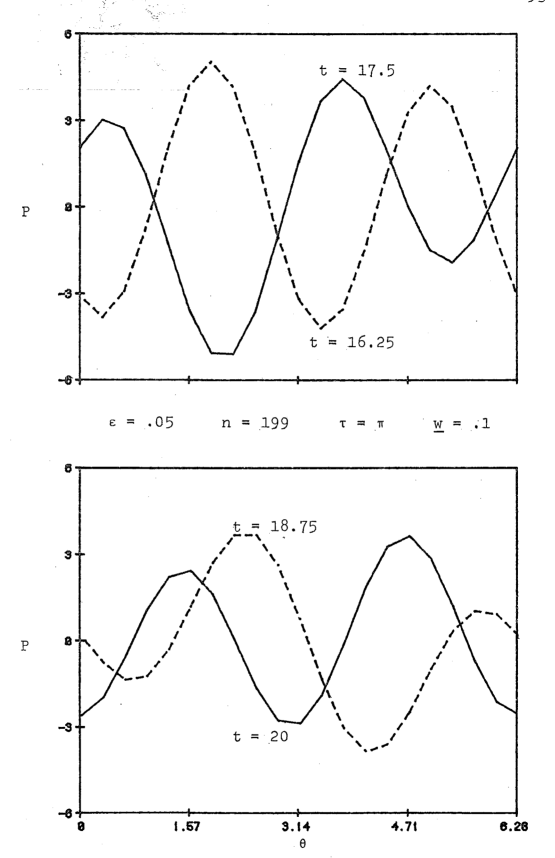


Figure 37. The Wall Pressure Waveforms for Traveling Waves

APPENDIX A

DERIVATION OF THE COEFFICIENTS

In this section, the coefficients b_{ij} , c_{ij} , and d_{ij} appearing in chapter 3 and 5 are derived and their numerical values are presented in Table 6, 7, and 8 respectively.

In chapter 3, for initial condition (3.7), one assumes

$$\partial_{tt}\phi_2 - \nabla^2\phi_2 = \sum_{m=0}^{\infty} H_m(r)\cos(m\theta)\sin(2S_{11}t).$$

By comparing coefficient yields

$$H_0(r) = \frac{1}{4}S_{11}(2S_{11}^2J_0^2 - (\gamma-1)S_{11}^2J_1^2 - 4S_{11}J_0J_1/r + 4J_0^2/r^2)$$

$$H_1(r) = 0$$

$$H_{2}(r) = \frac{1}{4}S_{11}(2S_{11}^{2}J_{0}^{2} - (\gamma-1)S_{11}^{2}J_{1}^{2} - 4S_{11}J_{0}J_{1}/r)$$

$$H_{m}(r) = 0$$
 for $m = 3, 4, ..., \infty$.

Then expanding ϕ_2 in a Fourier-Bessel series and expanding $H_\text{m}(\textbf{r})$ in a Bessel series leads to

$$b_{ij} = 2S_{ij}^{2} \int_{0}^{1} H_{i}(r) J_{i}(S_{ij}r) r dr / \{(S_{ij}^{2} - i^{2}) J_{i}^{2}(S_{ij})\}.$$

This integral must be computed numerically. For the sake of brevity only the first five coefficients in each series are calculated and presented in Table 6.

Table 6

BESSEL SERIES COEFFICIENTS FOR ANALYICAL SOLUTION

n	b _{0n}	b _{2n}
0 1 2 3 4 5	1.11738-0.37246 0.50462+0.37912 -0.09246-0.00802 0.05019+0.00184 -0.03290-0.00067 0.02375+0.00030	0.27181-1.094827 -0.11390-0.011107 0.05479+0.002087 -0.03451-0.000717 0.02451+0.000337

In Chapter 5, equation (5.8) can be written as

$$\phi = f_0 \phi_0 + f_1 \phi_1 + f_2 \phi_2 + g_1 \phi_3 + g_2 \phi_4 \tag{A-1}$$

where $\phi_0 = J_0$, $\phi_1 = J_1 \cos \theta$, $\phi_2 = J_2 \cos 2\theta$

$$\phi_3 = J_1 \sin \theta, \quad \phi_4 = J_2 \sin 2\theta.$$

By substituting (A-1) into the governing equation (5.6), yields

$$R = D(\phi) \tag{A-2}$$

where D is the nonlinear differential operator of equation (5.6). If (A-1) represented the exact solution to (5.6), R would vanish. Since this is not the case R has a finite value. The Galerkin procedure consists of making R orthogonal to each of the ϕ_1 's. This leads to the equations

$$\int_{0}^{1} \int_{0}^{2\pi} R\phi \operatorname{rdrd} = 0 \quad i = 0, 1, \dots, 4$$
 (A-3)

and a set of five second order differential equations are generated. The angular part of integration can be performed in closed form while the radial part of integration must be computed numerically and leads to the integral

$$C_{1} = \beta_{k} \int_{0}^{1} F_{1}(r) J_{k}(S_{k1}r) r dr \qquad (A-4)$$

$$\beta_{k} = 2S_{k1}^{2}/\{(S_{k1}^{2} - k^{2})J_{k}^{2}(S_{k1}^{2})\}. \tag{A-5}$$

The numerical values of C_i are listed in Table 7 for γ = 1.2. Also listed in this table are the functional forms of F_i 's. In writing these the notation

$$R_{i} = S_{i1}^{2}(\gamma-1)J_{i}^{"} + S_{i1}J_{i}^{!}/r - i^{2}J_{i}/r^{2}, \quad i=0,1,2$$
 (A-6)

is used. Similarly the numerical values of

$$d_{i,j} = \beta_i \int_0^1 G_{i,j} J_i(S_{i,1}r) r dr \qquad (A-7)$$

and functional forms of Gij's are listed in Table 8.

COEFFICIENTS APPEARING IN EQUATIONS (5.9) γ = 1.2

Table 7

i	k	c ₁	F _i
1.	0	4.137	R ₀ J ₀ + 2S ² ₁ J ²
2	0	1.042	$R_1 J_1/2 + S_{11}^2 J_1^{'2} + J_1^2/r^2$
3	0	-0.208	$R_2J_2/2 + S_{21}^2J_2^2 + 4J_2^2/r^2$
4	1	-1.939	$R_{0}J_{1} + 2S_{01}S_{11}J_{0}^{i}J_{1}^{i}$
5	1	-2.312	$R_{1}J_{0} + 2S_{01}S_{11}J_{0}^{i}J_{1}^{i}$
6	1	1.719	$R_1 J_2 / 2 + S_{11} S_{21} J_1 J_2 + 2 J_1 J_2 / r^2$
-7	1	1.483	$R_2J_1/2 + S_{11}S_{21}J_1'J_2' + 2J_1J_2/r^2$
8	2	-2.785	$R_0J_2 + 2S_{01}S_{21}J_0^{\dagger}J_2^{\dagger}$
9	2	-3.039	$R_{2}J_{0} + 2S_{01}S_{21}J_{0}^{\dagger}J_{2}^{\dagger}$
10	2	1.132	$R_{1}J_{1}/2 + S_{11}^{2}J_{1}^{2} - J_{1}^{2}/r^{2}$
11	0	2.586	$S_{01}^2J_0^2$
12	0	0.480	$\frac{1}{2}(S_{11}^2J_{1}^{12} + J_{1}^2/r^2)$
1,3	0	-0.196	$S_{21}^2 J_2^{12}/2 + 2J_2^2/r^2$
14	1	-4.849	2S ₀₁ S ₁₁ J'J' ₁
15	1	1.503	$S_{11}^{S_{11}^{J_1'J_1'} + 2J_1^{J_2/r^2}}$
16	2 .	-0.576	$\frac{1}{2}(S_{11}^2J_1^{2} - J_1^2/r^2)$
17	2	-3.481	2S ₀₁ S ₂₁ J'J' ₂

.

COEFFICIENTS APPEARING IN EQUATION (5.12)

Table 8

i	j	d ij	G i.j
0	1	1.000	J ₀ (S ₀₁ r)
0	2	1.293	¹ ₂ S ² ₀₁ J ¹ ₀ ²
0	3	0.240	$\frac{1}{4}(S_{11}^2J_1^2 + J_1^2/r^2)$
0	4	-0.098	$\frac{1}{2}(S_{21}^2J_2^{12}/2 + 2J_2^2/r^2)$
0	5	-0.176	-1 ₂ J ₀ ²
0	- 6	0.061	- ¹ 4J ²
0	7	0.049	- ¹ ₄ J ² ₂
1	1	1.000	J ₁ (S ₁₁ r)
1	2	-1.212	$S_{01}S_{11}J_0^{\dagger}J_1^{\dagger}$
1	3	0.927	$\frac{1}{2}(S_{11}S_{21}J_1^{\dagger}J_2^{\dagger} + 2J_1J_2/r^2)$
1	4	0.165	$-J_0J_1$
1	5	-0.199	-½J ₁ J ₂
2	1	1.000	J ₂ (S ₂₁ r)
2	2	-1.741	S ₀₁ S ₂₁ J'J' ₂
2	3	-0.223	$\frac{1}{4}(S_{11}^2J_1^{12} - J_1^2/r^2)$
2	4	0.237	$-J_0J_2$
2	5	-0.175	-¾J ² 1

APPENDIX B

PROGRAM FOR STABILITY BOUNDARY CURVES

```
SSET LIST FREE
C
      KIN-WING WONG
                                    JUN 30 1978
C
C
      CCMBUSTION INSTABILITY
C
      METHOD GALERKIN METHOD
Cź
      THIS PROGRAM GENERATE THE STABILITY CURVE.
C
      INPUT INFORMATION
C
C
      THE PROGRAM USED THE FORMAT FREE INPUT OUTPUT. A
C
      CCMMA SERVES AS A DELIMITER AND THE INPUT VARIABLES
C
      AFE ENTER IN THE FOLLOWING ORDER
         TI
                    INITIAL TIME
         TF
                    FINAL TIME
         DT
                    STEP SIZE FOR TIME
C
         WB
                    BURNING RATE
C
         TDELAY
                    DELAY TIME
C
C
         ΕI
                    INITIAL EPS
C
         EF
                    FINAL EPS
€
         DE
                    STEP SIZE FOR EPS
C
         DA
                    ASSUME POSITION FOR N
C
         FOFFD
                    ARE THE INITIAL CONDITION MODIFIER AND
C
                    HAVING THE VALUE EITHER 1 OR O
C
C
      ANOTHER SETS OF DATA CAN BE ENTER BY ENTERING
         EI, EF, DE, DA, FO AND FD.
C
      CIMENSION DY(10), FO(5), DF(5), G(10,500), Y(10)
      CCMMON EP,
                    RK. WB. EPANV
      CCMMON /SLIST/SO, S1, S2
      CCMMON /BLK1/C1,C2,C3,C4,C5,C6,C7,C8,C9,C10,C11,C12
     1, C13, C14,
                     C15,C16,C17
      LCGICAL PASS, GROW
      CATA MAXITR, EPS/20,-1/
      CATA PI/3.14159/
      READ(5,/) TI,TF,DT,WB,TDELAY
      PFINT*//, TI, TF, DT, WB, TDELAY
      TEELAY = PI * TDELAY
      K=TDELAY/DT
      K1=K+1
      IF (K1.GT.5CO) GO TO 105
      IF (K.LT.1.AND.NOF.EG.1) K=1
      NSTEP = (TF-TI)/DT
      h=10
      HEALF = DT * . 5
      NFROB=0
      READ(S,/,END=999) EI, EF, DE, DA, FO, CF
      NFROB=NPROB+1
```

```
FRINT *//, NPROB
       FFINT *//* FO.DF
       MSTEP=(EF-EI)/DE
       EF=EI
       IF (EP.LT.0) EP=.01
       ARV=DA
       FASS = . FALSE .
       CC 100 ISTEP =0 MSTEP
       CC 80 ITR=1, MAXITR
       EFAN V = EP * AN V
       IF (EPANV-EG-G) EPANY=ANV
       CC 10 I=1.K
       CC 10 J=1.N
 1 C
       G(J_PI)=0.
       CC 20 I=1.N
 20
       Y(I)=0.
C
C
       INITIAL CONDITION
C
       Y(2) = F0(1) \frac{1}{2}Y(4) = F0(2)\frac{1}{2}Y(6) = F0(3)\frac{1}{2}Y(8) = F0(4)\frac{1}{2}Y(10) = F0
      1(5)
       Y(1)=DF(1)*SO; Y(3)=DF(2)*S1; Y(5)=DF(3)*S2
       Y(7) = 0F(4) * S1; Y(9) = 0F(5) * S2
       CALL DLYFUN (NøYøGøKøEPSøK1)
       I=II
C
       CC 40 I=1, NSTEP
       CALL RKDELY (NoToYODYODTOGOKOEPO & 1100 HHALFOK1)
       CC 30 J=2.N.2
       1F(ABS(Y(J)).GT.2) GO TO 60
 3 C
       CENTINUE
       CENTINUE
 4 C
       GFOW= . FALSE .
       IF (PASS) GO TO 50
       AN VD WN = AN V
       ARV=ARV+DA
       GC TO 80
 50
       CENTINUE
       ANVDWA=ANV
       GC TO 70
       CCNTINUE
 60
       PASS=. TRUE.
       GFOW = . TRUE .
       ANVUP=ANV
 7 C
       CENTINUE
       IF (ABS(ANVUP-ANVDHN).LT.EPS) GO TO 90
       ANV= (ANVUP + ANVD WN) * . 5
 8 C
       CENTINUE
 90
       CENTIAUE
       NNU=ANV
       PRINT *//, EP, NNU, ITR
```

```
IF (ITR-EQ-MAXITR) PRINT //p* *** WARNING *** THE
                     1 ANSWERMAY NOT CONVERGE.
                        EP=EP+DE
                             ANV=ANVUP/23ANVDWN=0
    160
                        CENTINUE
                        GC TO 9
                        PFINT // DELAY TIME TOO LARGE " , TDELAY
   1 05
                         GC TO 999
    11C
                        PFINT *//, *ILLEGAL DELAY FUNCTION *
    999
                        SIOP
                        END
                         SUBROUTINE DIFFUN (T,Y,DY)
C
C
                        THIS SECTION PROVIDE A SET OF FIVE SECOND ORDER
C
                             EQUATIONS
C
                         WHERE
C
                                    F0=Y(2)
                                                                                                                       G1=Y(8)
C
                                    F1 = Y(4)
                                                                                                                       G2=Y(10)
C
                                    F2=Y(6)
                        DIMENSION Y(10), CY(10)
                        CCMMON EP. FK. WB. EPANV
                        COMMON /SLIST/SO,S1,S2
                        CCMMON /BLK1/ C1. C2. C3. C4. C5. C6. C7. C8. C9. C10. C11. C12
                    1, (13, (14,
                                                                             C15,C16,C17
C
                        [1(2)=y(1)]
                        DY(1) = -S0*S0*Y(2) - EP*(C1*Y(2)*Y(1)+C2*(Y(4)*Y(3)+Y(8)
                     1*Y(7)  *C3*(Y(10)*Y(9)*Y(6)*Y(5))
                        C1(4)=Y(3)
                        DY(3) = -S1 + S1 + Y(4) - EP + (C4 + Y(2) + Y(3) + C5 + Y(4) + Y(1) + 
                     1 C6*(Y(8)*Y(9)+Y44)*Y(5))+C7*(Y(6)*Y(3)*Y(10)*Y(7)))
                        CY(6)=Y(5)
                        C1(5)=-S2*S2*Y(6)~EP*(C8*Y(2)*Y(5)+C9*Y(6)*Y(1)+
                                                                                                                                                                                                                                       C10*
                     1(Y(8)*Y(7)=Y(4)*Y(3)))
                        E1(8)=Y(7)
                        EY(7)=-S1*S1*Y(8)-EP*(C4*Y(2)*Y(7)+C5*Y(8)*Y(1)*
                                                                                                                                                                                                                                       C6*
                     1(Y(4)*Y(9)-Y(8)*Y(5))+C7*(Y(10)*Y(3)-Y(6)*Y(7)))
                        0Y(10) = Y(9)
                        C1(9)=-S2*S2*Y(10)-EP*(C8*Y(2)*Y(9)+C9*Y(10)*Y(1)
                    1*(Y(8)*Y(3)*Y(4)*Y(7)))
                        IF (RK.EQ. 2) RETURN
C
C
                        THIS PROVIDE THE COMBUSTION TERMS
C
                        CY(1) = DY(1) - WB * (Y(1) + EPANV * (C11 * Y(2) * Y(2) + C12 * (Y(4) * Y(2) * Y(2) + C12 * (Y(4) * Y(2) 
                     1(4) + Y(8) + Y(8) +
                                                                                                               C13*(Y(10)*Y(10)+Y(6)*Y(6)))
                         Y(8)*Y(10))))
                         DY(5) = DY(5) - Wa*(Y(5) + EPANV*(C16*(Y(4)*Y(4)-Y(8)*Y(8))
                                                     +C17*Y(2)*Y(6)))
```

```
CY(7)=DY(7)-WB*(Y(7)+EPANV*(C14*Y(2)*Y(8)+C15*(Y(4)*Y
     1(10) Y(8) * Y(6)))
      DY(9)=DY(9)-W8*(Y(9)+EPANY*(C17*Y(2)*Y(10)+2*C16*Y(4)
     1 * Y(8)))
      RETURN
      END
      SUBROUTINE REDELY (NoTo Yo DY o Ho Go Ko EPS o * o HHALF o K 1)
C
С
      THIS SECTION PERFORM A FOURTH ORDER RUNGE-KUTTA METHOD
C
      WITH TIME DELAY FUNCTION
      CIMENSION Y(10), DY(10), Y2(10), Y3(10), G(10,500)
      CEMMON EP.
                    RK . WB . EPANV
      CCMMON /BLK1/ C1=C2=C3+C4+C5+C6+C7+C8+C9+C10+C11+C12
     1,C13,C14,
                    C15,C16,C17
      CALL DIFFUNCT, Y. DY)
      EC 10 I=1 N
          Y2(I)=Y(I) + HHALF*(DY(I) + G(I,1))
      G(I_{p}1)=(G(I_{p}1)+G(I_{p}2))/2.
 1 C
      CENTINUE
      T=T+HHALF
      CALL CIFFUN(T, Y2,DY)
      CC 20 I=1, N
          Y3(I)=Y(I) + HHALF*(DY(I) + G(I-1))
 20
          Y2(I) = Y2(I) + 2.*Y3(I)
      CALL DIFFUNC I, Y3, DY)
      CC 40 I=1.N
          Y3(I)=Y(I) + H*(DY(I) + G(I,1))
      IF(K.LT.1) GO TO 40
      CC 30 J=1.K
 3 C
      G(I \cap J) = G(I \cap J+1)
 4 C
          Y2(1) = Y2(1) * Y3(1)
      T=T + HHALF
      CALL DIFFUN (T ,Y3,DY)
      CC 50 I=1 N
          Y(I) = (Y2(I) - Y(I) + HHALF*(DY(I) + G(I+1)))/3.
 5 C
       CONTINUE
      ENTRY DLYFUN (N, Y, G, K, EPS, K1)
C
      THIS GIVES THE FORM OF DELAY FUNCTION G
C
      IF (K.LT.1) RETURN
      IF (RK.EQ.2) RETLAN
      G(1,K1) =
                  ₩B*
                              EPANV*(C11*Y(2)*Y(2)+C12*(Y(4)*Y
     1(4) + Y(8) * Y(8) +
                             C13*(Y(1G)*Y(10)*Y(6)*Y(6)))
      G(3,K1) =
                             EPANV*(C14*Y(2)*Y(4)+C15*(Y(4)*Y
                   ₩8*
             Y(8)*Y(10)))
     1(E)+
      G(5,K1)=WB*EPANV*(C16*(Y(4)*Y(4)*Y(8)*Y(8))+C17*Y(2)*Y(2)*Y(4)*Y(4)*Y(8)*Y(8)
     1(())
      G(7,K1)=
                   WB*
                             EPANV*(C14*Y(2)*Y(8)+C15*(Y(4)*Y
     1(10) Y(8)*Y(6))
```

```
G(9,K1)=WB*EPANV*(C17*Y(2)*Y(10)+2*C16*Y(4)*Y(8))
RETURN
END
BLOCK DATA
CGMMON /8LK1/C1,C2,C3,C4,C5,C6,C7,C8,C9,C10,C11,C12
1,C13,C14, C15,C16,C17
CEMMON /SLIST/S0,S1,S2
CATA S0,S1,S2/3.83171,1.84118,3.05424/
CATA C1,C2,C3,C4/4.1373,1.0423,-.2084,-1.9394/
CATA C5,C6,C7,C8,C9 /=2.3123,1.7187,1.4828,-2.785,
1-3.0388/
CATA C10,C11,C12,C13/1.1318,2.586,.480,-.196/
CATA C14,C15,C16,C17/-2.4243,1.8534445,-.447,-3.481/
END
```

APPENDIX C

PROGRAM FOR MODAL AMPLITUDE AND PRESSURE

```
SSET LISTUTOBIND
SBIND = FROM *FORTLIB/= FREE
      1(KINC=DISK, MAXRECSIZE=15, BLOCKSIZE=420, AREAS=4
     1 AREASIZE=1400 FILETYPE=7)
C
      KIN-WING WONG
C
                                    JUN 30 1978
C
      CCMBUSTION INSTABILITY
C
C
      METHOD: THIS PROGRAM USED THE GALERKIN METHOD TO
C
         GENERATE FIVE SECOND ORDER DIFFERENTIAL EQUATIONS.
C
          A FOURTH ORDER RUNGE-KUTTA METHOD IS USED TO SOLVE
C
         THE RESULTING EQUATIONS.
C
      THE PROGRAM GIVES THE MODAL AMPLITUDE AND THE PERTUBED
C
C
          PRESSURE IN GRAPHICAL FORM.
C
      THERE ARE THO TYPES OF INPUT DATAS. THE FIRST SET OF INPUT DATA USED FORMAT FREE INPUT. A COMMA SERVES
C
C
С
          AS A DELIMITER AND THE DATA ENTER AS FOLLOW:
C
               TI
                    INITIAL TIME
C
                    FINAL TIME
               TF
C
               DI
                    STEP SIZE FOR TIME
C
               NP
                    PRINT FREQUENCY
C
                    1 FOR THE DELAY TIME APPROCHING O
               NDF
C
                    O OTHERWISE
C
               IGAS 1 FOR GAS-DYNAMIC TERMS
C
                    O OTHERWISE
C
                    INITIAL CONDITION MODIFIER
               FO.
C
               DF
                    EITHER 1 OR 0
C
                    RADIUS OF THE CYLINDER
C
C
      THE SECOND SET OF DATA USED THE NAMELIST (LIST)
          INPUT OUTPUT OPTION. THE DATA CAN BE ENTERED IN
C
          ANY ORDER. HOWEVER THE FOLLOWING DATA MUST BE
C
          PROVIDED INITIALLY.
C
           TOELAY
                    DELAY TIME IN MULTIPLE OF PI
C
                    ORDER PARMETER
               EΡ
C
               мB
                    STEADY BURNING RATE
С
                    INTERACTION PARMETER
C
               PLOT T FOR PRESSURE PLOT
C
                    F OTHERWISE
C
      IF PLOT IS TRUE, ONE MORE DATA (NUMPT) IS NEEDED TO
C
          PROVIDE THE NUMBER OF POINT WITHIN O AND 2*PI.
      DIMENSION XX(100), YY(100), DY(10), P(5), FO(5), DF(5), G
     1(10,500),Y(10)
      CEMMON EP, IGAS,
                          RK, WB, EPANV
      CEMMON /SLIST/SO,S1,S2
      CCMMON /BLK1/ C1,C2,C3,C4,C5,C6,C7,C8,C9,C10,C11,C12
```

```
1,013,014, 015,016,017
      COMMON /BLK2/CO2,CO3,CO4,CO5,CO6,CO7,C12P,C13P,C14P
     1,C15P,
                          C22, C23, C24, C25, GAM A
      COMMON /BLK3/ BJO,BJ1,BJ2,COSQ,COS2Q,SINQ,SIN2Q
      DATA PI/3-14159/
      LCGICAL PLOT
      NAMELIST /LIST/ TDELAY, EP, WB, ANV, PLOT
      READ(5,/) TI,TF,DT,NP,NDF,IGAS,FO,DF,R
      PRINT *//, TI, TF, DT, NP, NDF
      IF (IGAS.EQ.O) PRINT //, GAS DYNAMIC TERM IS OFF
      PRINT *//,FO,DF,R
      EFS=1.E-6
      CALL BESJ(SO*R,O,BJO,EPS,IER)
      CALL BESJ(S1*R,1,BJ1,EPS,1ER)
      CALL BESJ(S2*R,2,BJ2,EPS,IER)
 999
      READ (5, LIST, END =99)
      REWIND 1
      TCLY=PI*TDELAY
      EFANV=EP*ANV
      IF (EPANV-EQ-0) EPANV=ANV
      WRITE(6,LIST)
      N=10
      HHALF=DT*.5
      K=TDLY/DT
      K1=K+1
      IF (K1 .GT. 500) GO TO 500
      IF (K.LT.1.AND.NDF.EQ.1) K=1
      DC 30 I=1.K
      DC 30 J=1.N
 30
      G(J,I) = 0.
      DO 10 I=1,N
 10
      Y(I)=0.
      INITIAL CONDITION
C
C
      Y(2) = F(0(1)) Y(4) = F(0(2)) Y(6) = F(3) Y(8) = F(4) Y(10) = F(4)
      Y(1) = DF(1) * SO; Y(3) = DF(2) * S1; Y(5) = DF(3) * S2
      Y(7) = DF(4) *S1; Y(9) = DF(5) *S2
      NSTEP=(TF-TI)/DT
      IF (.NOT.PLOT) GO TO 40
      CALL PRESUE(Y,P)
      WRITE(1) T>P
 4 C
      CENTINUE
      PRINT //, "FUNCTION"
      WRITE (6,100) T, (Y(1),1=2,N,2)
45
      CONTINUE .
      CALL DLYFUN (N,Y,G,K,EPS,K1)
      DC 20 I=1.NSTEP
      CALL RKDELY(N, T, Y, DY, DT, G, K, EP, &400, HHALF, K1)
      DC 55 J=2,N,2
```

```
IF(ABS(Y(J)).LT.2) GO TO 55
      RK=2
 55
      CONTINUE
      IF (MOD(I, NP).NE.O) GO TO 20
      IF (.NOT.PLOT) GO TO 60
      CALL PRESUE(Y,P)
      WRITE(1) T.P
 60
      ARITE(6,100) T_{*}(Y(J),J=2,N,2)
 65
      CONTINUE
 20
      CONTINUE
      IF (RK.GT.O) PRINT //, UNSTABLE
      IF (.NOT.PLOT) GO TO 70
      ENDFILE 1
      REWIND 1
      PRINT //, PRESSURE *
      READ(5,/) NUMPT
      DG=PI * 2./NUMPT
      NLM=NUMPT+1
      TEMP = GJXX(1) = OJ
      CC 75 I=2, NUM
      TEMP = TEMP + DQ
 75
      XX(I) = TEMP
 98
      READ(1, END = 70) T.P
      PRINT //>TIME = ">T
      SUMY = C.
      DC 80 I=1, NUM
      QG = XX(I)
      YY(I) = -GAMA*(P(1)*P(2)*COS(QQ)*P(3)*COS(2*QQ)*P(4)
                      P(5)*SIN(2*QQ))
     1*SIN(QQ)+
      SUMY=SUMY+YY(I)
      CONTINUE
 80
      IF (SUMY-NE-0.) CALL PLOTZD(XX,YY,NUM,100,100)
      GO TO 98
 70
      CONTINUE
      RK=0
      GG TO 999
 400
      PRINT *//, * ILLEGAL DELAY FUNCTION *
      GC TO 99
 50C
      PRINT *//, *DELAY TIME TOO LARGE *
 99
      SIDP
 100
      FERMAT (X, 8E12.5)
 110
      FORMAT(13X,7E12.5)
      END
      SUBROUTINE DIFFUN (T, Y, DY)
      CIMENSION Y(10), DY(10)
      COMMON EP, IGAS, RK, WB, EPANV
      COMMON /SLIST/SO,S1,S2
      COMMON /BLK1/ C1,C2,C3,C4,C5,C6,C7,C8,C9,C10,C11,C12
     1,C13,C14,
                    C15,C16,C17
C
      CY(2)=Y(1)
```

```
DY(1)==S0*S0*Y(2)=EP*IGAS*(C1*Y(2)*Y(1)+C2*(Y(4)*Y(3)
1 + Y(3) * Y(7)
              *C3*(Y(10)*Y(9)+Y(6)*Y(5)))
DY(4)=Y(3)
 DY(3) = -S1 * S1 * Y(4) - EP * IGAS * (C4 * Y(2) * Y(3) * C5 * Y(4) * Y(1) *
     C6*(Y(8)*Y(9)*Y(4)*Y(5))+C7*(Y(6)*Y(3)*Y(10)*Y
2(7)))
 DY(6) = Y(5)
DY(5)==S2*S2*Y(6)=EP*IGAS*(C8*Y(2)*Y(5)+C9*Y(6)*Y(1)+
1 C10*(Y(8)*Y(7)-Y(4)*Y(3)))
DY(8) = Y(7)
 DY(7)=-S1*S1*Y(8)-EP*IGAS*(C4*Y(2)*Y(7)+C5*Y(8)*Y(1)+
1 C6*(Y(4)*Y(9)=Y(8)*Y(5))+C7*(Y(10)*Y(3)=Y(6)*Y(7)))
DY(10) = Y(9)
DY(9)=-S2*S2*Y(10)-EP*IGAS*(C8*Y(2)*Y(9)+C9*Y(10)*Y(1)
   C10*(Y(8)*Y(3)*Y(4)*Y(7)))
 IF (RK.EQ. 2) RETURN
 DY(1)=DY(1)-W8*(Y(1)+EPANV*(C11*Y(2)*Y(2)+C12*(Y(4)*Y
       Y(8) *Y(8)) *
                      C13*(Y(10)*Y(10)*Y(6)*Y(6))))
 CY(3)=DY(3)-WB+(Y(3)+EPANV+(C14+Y(2)+Y(4)+C15+(Y(4)+Y
1(6)+
        Y(8) = Y(10)))
 CY(5) = DY(5) = WB * (Y(5) * EP ANV * (C 16 * (Y(4) * Y(4) = Y(8) * Y(8))
        +C17*Y(2)*Y(6)))
 1(10) \quad Y(8) * Y(6)))
 CY(9) = DY(9) - WB* (Y(9) + EPANV* (C17*Y(2)*Y(10) + 2*C16*Y(4))
1 * Y(8)))
RETURN
END
 SUBROUTINE RKDELY(N, T, Y, DY, H, G, K, EPS, *, HHALF, K1)
 DIMENSION Y(10), DY(10), Y2(10), Y3(10), G(10,500)
COMMON EP, IGAS,
                    RK, WB, EPANV
 CCMMON /BLK1/ C1,C2,C3,C4,C5,C6,C7,C8,C9,C10,C11,C12
1, C13, C14,
              C15,C16,C17
CALL DIFFUNCT, Y. DY)
 00 1 I=1.N
    Y2(I)=Y(I) + HHALF*(DY(I) + G(I,1))
G(I+1)=(G(I+1)+G(I+2))/2
 CONTINUE
 T=T+HHALF
CALL DIFFUN(T, Y2, DY)
 00 2 I=1.N
    Y3(I)=Y(I) + HHALF*(DY(I) + G(I,1))
    Y2(I) = Y2(I) + 2*Y3(I)
 CALL DIFFUN( T, Y3, DY)
 DO 3 I=1.N
    Y3(I)=Y(I) + H*(DY(I) + G(I=1))
 IF(K-LT-1) GO TO 3
 DC 10 J=1,K
 G(I_{\rho}J)=G(I_{\rho}J+1)
    Y2(1)=Y2(1) + Y3(1)
T=T+HHALF
```

1

2

10

3

```
CALL DIFFUN (T , Y3, DY)
     DC 4 I=1.N
        Y(I) = (Y2(I) - Y(I) + HHALF*(DY(I) + G(I,1)))/3.
     CONTINUE
4
     FNTRY
            DLYFUN (NaYaGaKaEPSaK1)
     IF (K.LT.1) RETURN
     IF (RK.EQ.2) RETURN
     G(1)Ki)=
                 MB*
                           EPANV*(C11*Y(2)*Y(2)*C12*(Y(4)*Y
    1(4)+
          Y(8) *Y(8))+
                           C13*(Y(10)*Y(10)*Y(6)*Y(6)))
     G(3,K1) =
                           EPANV*(C14*Y(2)*Y(4)*C15*(Y(4)*Y
                 WB *
    1(6)+
            Y(8)*Y(10)))
     G(5,K1)=WB*EPANV*(C16*(Y(4)*Y(4)-Y(8)*Y(8))+C17*Y(2)*Y
    1(6))
     G(7 \cdot K1) =
                  WB*
                           EPANV*(C14*Y(2)*Y(8)+C15*(Y(4)*Y
          Y(8) *Y(6)))
     G(9,K1)=WB*EPANV*(C17*Y(2)*Y(10)+2*C16*Y(4)*Y(8))
     RETURN
     END
     SUBROUTINE PRESUE(Y,P)
     THIS SECTION PERFORM THE PRESSURE CALCULATION
     DIMENSION Y(10),P(5)
     COMMON EP, IGAS,
                        RK > WB > EPANV
     CCMMON /BLK2/CO2,CO3,CO4,CO5,CO6,CO7,C12P,C13P,C14P
    1,C15P,
                        C22, C23, C24, C25, GAMA
     CCMMON /BLK3/BJ0,BJ1,BJ2,COS0,COS20,SINQ,SIN20
     DF0=Y(1);DF1=Y(3);DF2=Y(5);DG1=Y(7);DG2=Y(9)
      F0=Y(2); F1=Y(4); F2=Y(6); G1=Y(8); G2=Y(10)
     P(1)=8J0*(DF0*EP*(C02*F0*F0+C03*(F1*F1+G1*G1)+C04*(F2
                     +C05*DF0*DF0+C06*(DF1*DF1+DG1*DG1)+C07*
    1*F2+G2*G2)
    2(DF2*DF2+
                    DG2*DG2)))
     P(2)=8J1*(DF1+EP*(C12P*F0*F1+C13P*(F1*F2+G1*G2)+C14P
    1 * 0 F 0 * 0 F 1 +
                    C15P*(DF1*DF2+DG1*DG2)))
     P(3) = 8J2*(DF2+EP*(C22*F0*F2+C23*(F1*F1=G1*G1)+C24*DF0)
    1*0F2+
                 C25*(DF1*DF1-DG1*DG1)))
     P(4) = BJ1 * (DG1 * EP * (C12P * F0 * G1 + C13P * (F1 * G2 = G1 * F2) * C14P
    1 * CF O * DG 1 +
                     C15P*(DF1*DG2-DF2*DG1)))
     P(5)=BJ2*(DG2*EP*(C22*F0*G2*2.*C23*F1*G1*C24*DF0*DG2*
          2.*C25*DF1*DG1))
     RETURN
     END
     BLOCK DATA
     CCMMON /BLK1/ C1,C2,C3,C4,C5,C6,C7,C8,C9,C10,C11,C12
    1,013,014,
                   C15,C16,C17
     COMMON /SLIST/SO,S1,S2
     CEMMON /BLK2/CO2,CO3,CO4,CO5,CO6,CO7,C12P,C13P,C14P
    1,015P,
                        C22, C23, C24, C25, GAMA
     DATA SO,S1,S2/3.83171,1.84118,3.05424/
     DATA C1, C2, C3, C4/4.1373, 1.0423, -. 2084, -1.9394/
     DATA C5,C6,C7,C8,C9 /-2.3123,1.7187,1.4828,-2.785,
```

C

C

1-3.0388/
DATA C10,C11,C12,C13/1.1318,2.586,.480,-.196/
DATA C14,C15,C16,C17/-2.4243,1.8534445,-.447,-3.481/
DATA C02,C03,C04,C05,C06,C07/1.2930,0.2400,-.0982,
1-.1762 ,.0607,.0494/C12P,C13P,C14P,C15P/-1.2121
2,.9267,.1651,-.1987/C22,C23,C24,C25/-1.7406,-.2235
3,.2371,-.1754/
DATA GAMA/1.2/
END

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16. Abstract									
The primary objective of this study is the development of a new analytical technique to be used in the solution of nonlinear velocity-sensitive combustion instability problems. Such a method should be relatively easy to apply and should require relatively little computation time. In an attempt to achieve this aim, the orthogonal collocation method is investigated first. However, it is found that the results are heavily dependent on the location of the collocation points and characteristics of the equations. Therefore, the method is rejected as unreliable. Next, the Galerkin method, which has proved to be very successful in analysis of the pressure sensitive combustion instability, is considered. This method is found to work very well. It is found that the pressure wave forms exhibit a strong second harmonic distortion and a variety of behaviors are possible depending on the nature of the combustion process and the parametric values involved. Finally, a one-dimensional model provides further insight into the problem by allowing a comparison of Galerkin solutions with more exact finite-difference computations.									
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