

11  
DOT/FAA/AM-94/X

Office of Aviation Medicine  
Washington, D.C. 20591

The Potential for Pulmonary Heat  
Injury Resulting from the Activation  
of a Cabin Water Spray System to  
Fight Aircraft Cabin Fires

Robert P. Garner, Ph.D.

Civil Aeromedical Institute  
Federal Aviation Administration  
Oklahoma City, OK 73125

This document is available to the  
public through the National Technical  
Information Service, Springfield,  
Virginia, 22161

TECHNICAL REPORT DOCUMENTATION PAGE

1. Report No. DOT/FAA/AM-94/X
4. Title and Subtitle  
The Potential for Pulmonary Heat Injury Resulting from  
the Activation of a Cabin Water Spray System to Fight  
Aircraft Cabin Fires
5. Report Date May 1994
7. Author(s)  
Robert P. Garner, Ph.D.
9. Performing Organization Name and Address  
FAA Civil Aeromedical Institute  
P.O. Box 25082  
Oklahoma City, OK 73125
12. Sponsoring Agency Name and Address  
Office of Aviation Medicine  
Federal Aviation Administration  
800 Independence Avenue, S.W.  
Washington, D.C. 20591
15. Supplementary Notes
16. Abstract

A cabin water spray system (CWSS) has been suggested as a means of attenuating the severity of smoke and fire commonly associated with aircraft accidents. All aspects of passenger and cabin safety must be considered when evaluating a new safety system or concept. The purposes of this report are to briefly review the pathophysiological changes occurring in the respiratory system as a result of thermal injury and to quantitatively estimate the risk of creating a more hazardous cabin environment by activation of a CWSS. Changes in the heat content of the cabin atmosphere resulting from CWSS activation were calculated using parameters consistent with current aircraft and proposals for CWSS design. The results suggest that only a very small volume of the aircraft cabin would have an increase in heat content that could result in thermal injury.

17. Keywords  
aviation; thermal injury; respiratory system; cabin water  
spray system

18. Distribution Statement

Document is available to the public through the  
National Technical Information Service, Springfield,  
Virginia 22161.

19. Security Classification

Unclassified

20. No of pages

16

## **I. Introduction**

Fire represents one of the most catastrophic events to which an organism can be subjected. Injury and death from fire can occur due to the incineration of tissue and also by the incapacitating byproducts of the fire environment. The odds of fire injury are increased when individuals are trapped within an enclosed space, such as an aircraft cabin. Testimony to this problem aboard aircraft is the Manchester incident of 1982, in which 52 people lost their lives due to inhalation injury. Efforts continue to develop safety systems that will minimize passenger deaths and injuries from aircraft fires. To date, no acceptable solution has been obtained. One suggestion has been to install some type of cabin water spray system (CWSS) in passenger aircraft to slow the spread of fire, thereby allowing more time for passenger evacuation. Because of the cost and complexity of such systems, implementation must be carefully evaluated.

One consideration of CWSS development is its impact on the fire environment. The use of a CWSS might result in the production of steam. Steam may have untoward effects on individuals in the area in which the CWSS was activated, thereby preventing any net benefits on passenger survival. The primary purposes of this paper are to examine the likelihood of 1) the water released from a CWSS being vaporized, to some extent, and causing respiratory tract injury, and 2) increasing the risk of thermal injury by temperature changes in the environment due to such vaporization. Some of this analysis will be based on experimental and clinical studies dealing with thermal injury, and some of it on theoretical calculations of heat content within the cabin environment in which a CWSS is heat activated.

## II. Mechanisms of Pulmonary Injury Due to Fire

Clinically, there are three categories of burn injury: (1) smoke inhalation, (2) cutaneous burn without smoke inhalation and (3) a combination of 1 & 2 (19). Pulmonary damage due to smoke inhalation and carbon monoxide intoxication is thought to be the leading cause of death in fire victims. Of 12,000 annual fire deaths in the United States, 50 to 60 per cent are secondary to inhalation injury (29). Careful study of smoke inhalation and the recognition of its existence did not begin until after the Coconut Grove night club fire, which occurred in Massachusetts in 1942. Thirty-six of the 39 individuals transported to the hospital eventually died of pulmonary injury, rather than cutaneous burns (2). Concern over this and other fire-related catastrophes developed into a significant interest in burn injury during the early 1940s. Investigation of the mechanisms of pulmonary injury, pathophysiological characteristics of the thermally injured respiratory tract, and optimization of diagnostic and treatment protocols has received worldwide investigative attention in the ensuing decades.

A fire represents a variety of chemical reactions, each of which contributes different groups of potentially injurious substances to the breathable atmosphere (28). The nature and amount of these combustion products are determined by the material being burned, the amount of oxygen that is available at different times during the conflagration, and the temperatures attained (18). Because of the fire environment's great complexity (25), it is difficult to make a definitive, quantitative assessment of those components of the fire atmosphere most responsible for inhalation injury. Direct injury of the respiratory system may be produced by inhalation of thermal, gaseous, and/or particulate matter. Inhalation injury resulting from fire is found most often with a combination of three conditions: (1) a closed-space accident, (2) the presence of

heavy smoke, and (3) a history of unconsciousness (20). Table 1 lists favorable and unfavorable conditions influencing the development of inhalation injury. Note that many of the factors that favor the development of the injury could be expected to be present during an aircraft cabin fire.

Factor	Unfavorable	Favorable
Place burn occurs	Enclosed (room, building, automobile)	Open
Anatomical location of burns	"Respiratory area" involved (mouth, pharynx, nose, nasal hairs)	"Respiratory area" spared
Type of burn	Flame	Liquid, chemical, irradiation
Heavy smoke; products of incomplete combustion	Present	Absent
Steam; high humidity	Present	Absent

Table 1. Factors influencing the incidence and severity of respiratory burns (23).

Victims of conflagrations frequently sustain more severe life-threatening pulmonary injuries than the burns received on the surface of the body. Wound sepsis was once the primary cause of death in burn patients. Since wound sepsis can now be controlled, respiratory complications of inhalation injury and pulmonary sepsis are currently a major cause of death in patients with body surface burns (11, 15). The incidence of significant pulmonary injury has been estimated to be between 15 and 22 percent in burn patient populations (1, 26) with inhalation injuries occurring in approximately one-third of all major burns (11).

Upper airway and lung inhalation injuries are usually secondary to inhalation of gaseous or particulate products of incomplete combustion and rarely are due to heat, unless steam has been inhaled. In some instances, respiratory tract damage is confined to the upper air passages

with little or no damage to the lungs. It should also be noted that death from inhalation injury can occur without thermal wounds. However, respiratory tract damage is often more severe in burned than in nonburned patients for a given degree of smoke inhalation. Death from inhalation injury is particularly common during the early postburn period (9, 19). The role that thermal, gaseous, and particulate matter may play in the development of respiratory damage from fire exposure is presented below.

### *Thermal Injury*

The definitive experimental study addressing the question of mechanism of thermal injury to the respiratory tract during a fire was published in 1945 (18). In that experiment, dogs inhaled hot air (350°C and 500°C), flame from a burner, and live steam. Only the steam resulted in thermal injury at the level of the lung parenchyma. The difference between the hot air and steam treatments were attributed to the low heat-carrying capacity of air coupled with the efficient heat absorption of the upper airway. Based on these observations, true thermal injury to the lower respiratory tract should be relatively rare. This is substantiated clinically by the observation that inhalation injury below the level of the vocal cords is a relatively uncommon consequence of thermal trauma, occurring in <5% of all burn patients (6, 22, 24). However, thermal injury to the respiratory tract is important. The unique problems of respiratory tract damage resulting from thermal injury have been recognized throughout the world (14, 27, 30). It has been suggested that classification of pulmonary burns be based on the etiology and severity of injury (4) instead of the time of clinical onset of respiratory distress, anatomy, or burn etiology.

### ***Particulate Injury***

Except for the rare instance of steam inhalation, direct thermal burns to the respiratory tract below the level of the larynx do not occur in patients (5, 6, 22, 24). The clinical "respiratory burn" is actually a chemical burn, induced by inhalation of suspended particles (smoke) and the toxic products of incomplete combustion (3, 8). This type of injury is a major complication of fire accidents. Chemical burn of the airways occurs upon inhaling the incomplete products of combustion, causing surface damage to the larynx, proximal and distal airways, and the lung parenchyma (13). The noxious constituents of smoke are believed to stimulate irritant receptors, producing bronchoconstriction and chemical injury to the airway mucosa and the alveolar-capillary membrane, thereby producing pulmonary edema. The interaction and impact of gases and other fire byproducts upon physiological function is extensively reviewed elsewhere (16, 25). The probability of respiratory tract injury resulting from inhalation of noxious gases and chemicals is much greater than thermal damage occurring from exposure, either to hot air saturated with water vapor, or live steam produced by the vaporization of droplets, which could result from use of a CWSS. Some of the morphological changes that could be expected from this type injury are presented below.

### **III. Pathophysiological Response to Thermal Injury**

Experimentally, evaluation of thermal inhalation injury in dogs has shown that inhalation of hot air (350 and 500°C) causes a thermal tracheitis of the upper trachea without injury to the lower trachea. Inhalation of steam causes thermal injury extending from the trachea to the lung parenchyma (18). Several breaths of steam delivered into the pharynx at a temperature of 100°C



causes such a severe local edema that the animals die within a few hours of obstructive asphyxia. Steam produces an immediate pulmonary edema in isolated lung lobes and the fluid transudation reaches its peak within half-an-hour after the inhalation (31). The ulcerations and rapid onset of edema formation observed in these studies is affirmed by clinical observations of inhalation injury.

Inhalation injury has been anatomically divided into three levels: 1) upper airway injury (burn damage limited to the larynx and vocal cords); 2) major airway injury (burn damage involving the tracheobronchial tree) and 3) parenchymal injury (burn damage involving the terminal bronchi and alveolar space) (21). Although the mucous membrane of the entire respiratory tract is potentially susceptible to damage, the infraglottic airway is often spared or injured to a lesser degree than the supraglottic airway because the vocal cords represent an anatomic barrier to the passage of heat into the trachea (13). This factor contributes to the relative rarity of true heat inhalation injury.

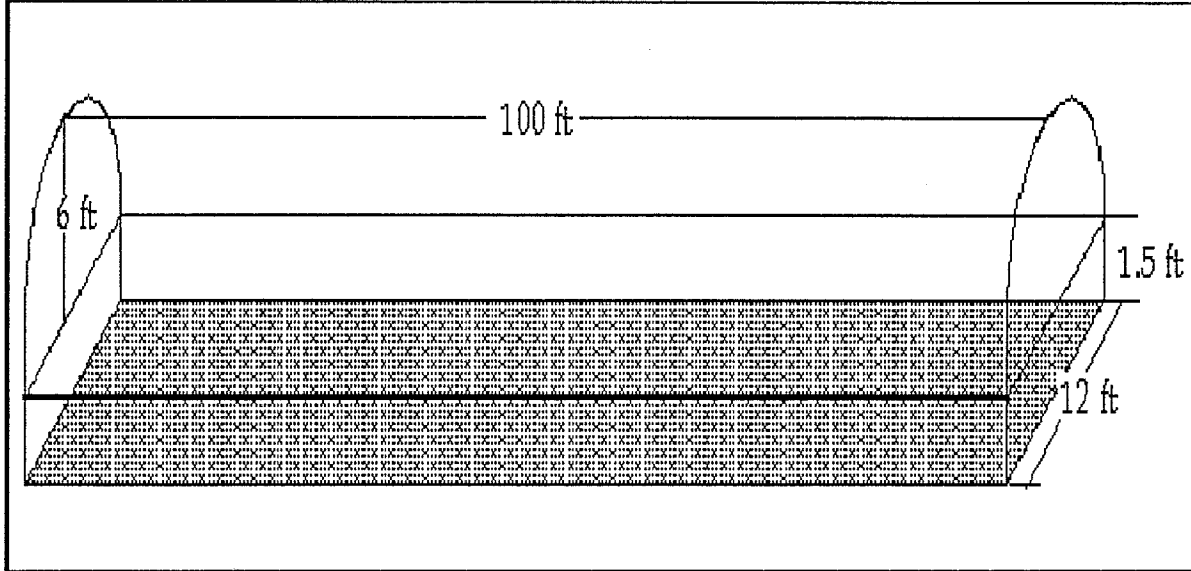
Functional and anatomical changes resulting from thermal injury to the respiratory tract have been well documented (7, 10). Heat energy produces immediate injury to the mucosa of the airway and pulmonary parenchyma. There is a rapid loss of fluid volume from the vascular space and concomitant expansion of the interstitial space (edema) immediately following injury. The mucosa develops severe edema, hemorrhage, and ulceration soon after exposure to hot steam. The parenchyma exhibits marginal emphysema and congestion, leading to acute pulmonary edema. Edema formation and congestion occur more rapidly in steam inhalation injury than in inhalation injuries produced by chemical agents (26). The tracheal lumen accumulates epithelial cells, white blood cells, and mucus, followed by the development of a

purulent and necrotizing broncho-pneumonia within 3 to 4 days.

Three major phases of respiratory injury induced by steam burns have been defined (21). The first phase (within the first hour) is identified by coagulation necrosis and an early reactive stage, with edema in the tracheobronchial tree and early pulmonary parenchymal edema. During the second phase (up to 24 hours), there is a second reactive stage, with development of interstitial and perivascular edema, further sloughing of mucosa, atelectasis, and hemorrhagic consolidation. Laryngeal edema and laryngospasm, leading to extrinsic obstruction of the swollen tissues around the vocal cords often occurs within the first 24 hours (11). The third, or infection phase (> 24 hours), is indicated by the development of bronchopneumonia behind respiratory tract obstruction, secondary to mechanical or functional block. Necrotizing tracheobronchitis also occurs in this phase.

#### **IV. Injury and Incapacitation due to Enhanced Heat Content of the Environment**

In assessing the toxicity of combustion products in fires, Purser (6) lists heat stroke, body surface burns, and respiratory tract burns as ways in which heat may lead to incapacitation and death in fire victims. The mechanisms and results of respiratory tract burns are discussed in Sections II and III above. With regard to CWSS activation, it is assumed that most individuals are evacuating and/or removed from areas in which the flames could result in direct injury. Therefore, potential body surface or respiratory tract burns would have to result from increased heat content of the environment. Based upon both clinical and experimental research observations, the only way in which heat content could have a significant impact is through the spray of the CWSS being converted to steam, or the heated air of the fire environment becoming



**Figure 1** Dimensions used for calculating cabin volume. A bisected cylinder and rectangular shapes were used for volume estimates.

saturated with this spray. What is the threat of thermal injury resulting from this scenario?

Each incident in which a CWSS is heat activated may be characteristically different. Prediction of potential water spray transformations is difficult. However, within the context of thermal injury, some basic assumptions must be made. Full-scale fire testing indicates that the presence of a CWSS lowers the rate of temperature increase under a variety of spray discharge configurations (17). Without water spray, cabin temperature underwent a drastic rise, starting at approximately 90 seconds, with temperature values of approximately 275°F being obtained at test termination time of 5 minutes. The current hypothesis holds that optimal CWSS performance requires the use of a zoned system in which water is discharged when air temperatures reach 300°F in a given zone (31). Based on this information, a temperature level of

300°F seems reasonable for use in calculations. For the entire activation and discharge period, a system using between 8 and 20 gallons of water is under consideration. Since one goal to be engineered into the final design of the CWSS is to minimize the volume of water necessary for effective operation, a volume of 10 gallons will be used for estimation. Cabin volume is estimated to be 7500 ft<sup>3</sup> (Figure 1).

The worst-case scenario is that the entire 10 gallons of water would be immediately converted to steam while an individual was located within the area of the steam cloud. If the initial temperature of the water in the CWSS was 75°F and the entire 10 gallons in the system were converted to steam at 120°F, ~24,000 kcal would be required. If this heat were supplied by dry air at 300°F, ~10,000 ft<sup>3</sup> of air would be required. However, based on cabin dimensions given above, only 75 ft<sup>3</sup> of air would be present at the hot-air/water-spray interface. This volume of air contains enough heat to convert ~0.076 gallons of water to steam, which represents a volume of 0.84 ft<sup>3</sup>. Therefore, it appears that the immediate formation of a significant steam cloud is unlikely.

The more likely occurrence would be that, as the water is discharged from the system, it saturates the heated air and this hot air and water vapor mixture diffuses "downstream" of the fire. This mixture presents a considerable threat of thermal injury due to the high heat content of water vapor and the potential condensation of this vapor in the lung. Table 2 lists the total heat content of water-saturated air in kcal/min and watts, based on a ventilatory rate of 10 l/min.

The heat delivered to the lung from inhaling dry or moist air of any temperature is found by subtracting the heat present in the ventilation volume from the heat contained in the air mixture of interest. From the inhaled air mixture temperature, the mean specific heat for both

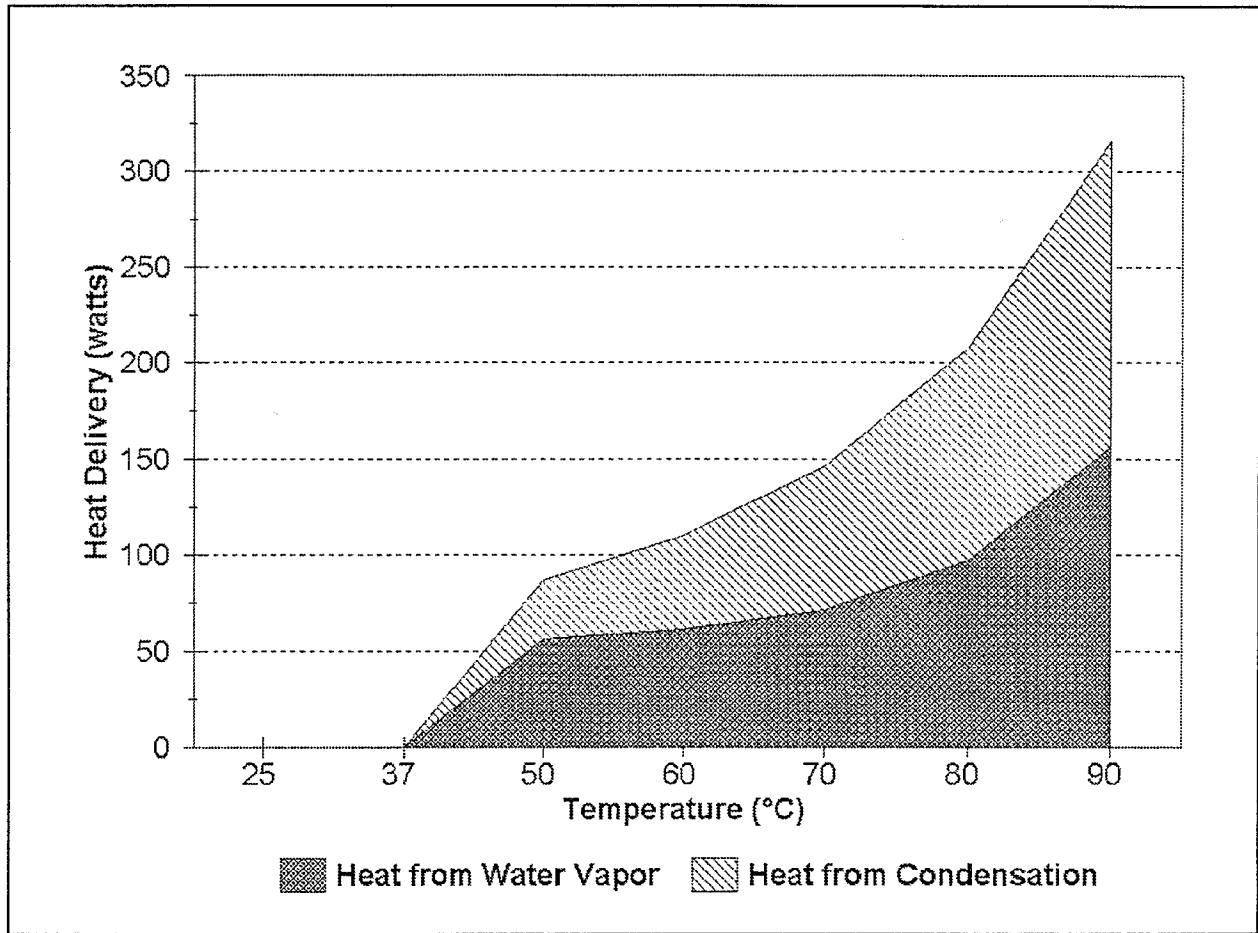
the water and air components can be determined. The density and partial pressure of each component must also be known. From these values, the heat per unit volume for each component in the mixture can be calculated. If a saturated water vapor mixture were inhaled at a temperature above 98.6°F, the heat of condensation of the water saturated air must be considered. The heat of condensation represents the majority of heat delivered by a water saturated air mixture.

Temperature (°F)	Temperature (°C)	kcal/min	Watts
122	50	1.253	87.42
140	60	1.583	110.49
158	70	2.091	145.94
176	80	2.973	207.44
194	90	4.514	315.09

Table 2. Total heat delivery to the lungs from breathing water vapor saturated air at various temperatures.

Figure 2 illustrates the quantity of heat that would be delivered to the lungs, based on a ventilatory rate of 10 l/min. As can be seen from the graph, an increasingly larger percentage of heat comes from the condensation of water vapor within the lung as the temperature increases. Experiments studying the impact of inhalation of hot water vapor saturated air have not been done. The expected response to this type of exposure would be reflex closure of the glottis which is the typical response to hot air (13). Therefore, significant damage in the areas of gas exchange would probably not occur. Of course, damage to the skin, nasal passages, and airways above the level of the glottis may occur. The extent to which this damage may occur would be dictated by the combination of temperature and exposure time. Potentially, exposure time would

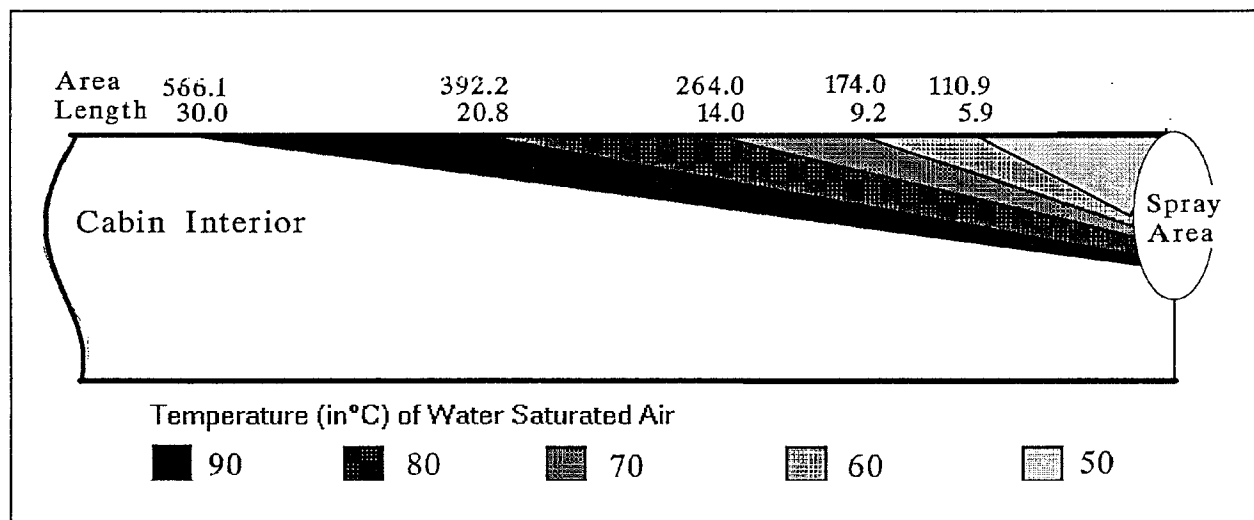
be correlated with the volume of saturated air present. This is particularly true if individuals are delayed in evacuating the aircraft cabin. What is the volume that poses this threat if 10 gallons of water are used to produce saturated air at the temperatures listed in Table 2?



**Figure 2** Heat delivery to the lungs based on inhalation of air saturated with water vapor.

Figure 3 presents an analysis for 10 gallons of water being used to create a saturated volume of air at the temperatures listed in Table 2. It is assumed that the saturated air mixture takes the form of a bisected cone as it moves away from the spray area along the ceiling of the aircraft cabin. As one would expect, the cooler saturated air is less of a threat. This is caused by: 1) a lower heat capacity, and 2) a reduced volume due to a lower density, as can be seen in

Figures 2 and 3, respectively. As temperatures increase, the threat of thermal injury becomes more pronounced. If someone were trapped in a saturated air environment above 70°C for more than a few seconds, it is very likely that serious injury or death would occur. However, it should be noted that the volume of the hottest saturated air represents only 7.6% of the total cabin volume. Furthermore, if the assumption that the high heat area is distributed along the top of the cabin, as depicted in Figure 3, much of the high heat exposure could be avoided by crouching or crawling near the floor during evacuation.



**Figure 3** Theoretical distribution of water saturated air in the aircraft cabin.

### *Conclusions*

Based on the worst-case analysis presented above, the risk due to increased latent heat in the environment resulting from activation of a CWSS is relatively small. Although a potential hazard from steam and hot water vapor saturated air does exist, exposure to these conditions for more than a second or two is highly unlikely, and could theoretically be avoided by maintaining the correct posture and quickly evacuating the aircraft. It should be noted that if the CWSS were

not present, the heat content of the environment would be significantly higher. The water of the spray absorbs heat which is, in essence, what produces a significantly lower rate of temperature rise in the cabin when a CWSS is activated (30). The fact that a relatively small, potentially hazardous thermal environment may be produced seems inconsequential, in comparison to an uncontrolled fire environment.



## References

1. Achauer, B.M., Allyn, P.A., Furnas, D.W. & Bartlett, R.H. Pulmonary complications of burns: the major threat to the burn patient. *Ann. Surg.*, 17: 311-319, 1973.
2. Aub, J.C., Pittman, H. & Brues, A.M. The management of the Cocoanut Grove burns at Massachusetts General Hospital: the pulmonary complications: a clinical description. *Ann. Surg.*, 117: 834-840, 1943.
3. Cohn, A.M. Concepts in management of burns of the respiratory tract. *Southern Med. J.*, 66: 297-301, 1973.
4. Chu, C.S. New concepts of pulmonary burn injury. *J. Trauma*, 21(11): 958-961, 1981.
5. Chu, C.S. Burns updated in China: II. Special burn injury and burns of special areas. *J. Trauma*, 22(7): 574-580, 1981.
6. DiVincenti, F.C., Pruitt, B.A., & Reckler, J.M. Inhalation injuries. *J. Trauma*, 11(2): 109-117, 1971.
7. Haponik, E.F., Meyers, D.A., Munster, A.M., Smith, P.L., Britt, E.J., Wise, R.A. & Bleeker, E.R. Acute upper airway injury in burn patients. *Am. Rev. Resp. Dis.*, 135: 360-366, 1987.
8. Harrison, H.N. Respiratory tract injury, pathophysiology and response to therapy among burned patients. *Ann. NY Acad. Sci.*, 150: 627-638, 1968.
9. Harvey, J.S., Watkins, G.S., & Sherman, R.T. Emergency burn care. *Southern Med. J.*, 77: 204-214, 1984.
10. Head, J.M. Inhalation injury in burns. *Am. J. Surg.*, 139: 508-512, 1980.
11. Heimbach, D.M. & Waeckerle, J.F. Inhalation Injuries. *Annals of Emerg. Med.*, 17(12): 1316-1320, 1988.
12. Hill, R.G., Marker, T.R., & Sarkos, C.S. Evaluation and optimization of an on-board water spray fire suppression system in aircraft. Paper presented at *12th Meeting of United States - Japan Panel of Fire Research and Safety*, Tsukuba and Tokyo, Japan, 1992.
13. Hunt, J.L., Agee, R.N., & Pruitt, B.A. Fiberoptic bronchoscopy in acute inhalation injury. *J. Trauma*, 15(8): 641-649, 1975.

14. Kerlakin, L.M. The thermal burn of the respiratory tract. *Vestnik Khirurgii*. 11: 41-47, 1962. Russian.
15. Loke, J., Matthay, R.A. & Walker Smith, G.J. The toxic environment and its medical implications with special emphasis on smoke inhalation. In Loke, J.(ed), *Pathophysiology and Treatment of Inhalation Injuries*, New York, Marcel Dekker, 1988.
16. Man, S.F.P. & Hulbert, W.C. Airway repair and adaptation to inhalation injury. In Loke, J.(ed), *Pathophysiology and Treatment of Inhalation Injuries*, New York, Marcel Dekker, 1988.
17. Marker, T. Onboard cabin water spray system under various discharge configurations. DOT/FAA/CT-TN91/42, 1991.
18. Moritz, A.R., Henriques, F.C., & McLean, R. The effects of inhaled heat on the air passages and lungs. *Am. J. Pathol.*, 21:311-331, 1945.
19. Mosley, S. Inhalation injury: A review of the literature. *Heart & Lung*, 17(1):3-9, 1988.
20. Moylan, J.A. Inhalation injury. *J. Trauma*, 21: 720-721, 1981.
21. Moylan, J.A. & Chan, C.K. Inhalation injury - an increasing problem. *Ann. Surg.*, 188(1): 34-37, 1978.
22. Phillips, A.W., Tanner, J.W., & Cope, O. Burn therapy. IV. Respiratory tract damage (an account of the clinical, X-ray and postmortem findings) and the meaning of restlessness. *Ann. Surg.*, 158: 799-811, 1963.
23. Pierson, D.J. Respiratory complications in the burned patient: pathophysiology and management. *Respiratory Care.*, 21(2): 123-133, 1976.
24. Pruitt, B.A., Erickson, D.R., & Morris, A. Progressive pulmonary insufficiency and other pulmonary complication of thermal injury. *J. Trauma*, 15: 369-379, 1975
25. Purser, D.A. Toxicity assessment of combustion products. in *SPFE Handbook of Fire Protection Engineering*, Society of Fire Protection Engineers, Boston, 1988.
26. Stone, H.H. & Martin, J.D. Jr. Pulmonary injury associated with thermal burns. *Surg. Gynec. Obstet.*, 129: 1242-1246, 1969.
27. Sheu, Y.H. The thermal burn of the respiratory tract. *Chinese Surg. J.*, 12: 762-770, 1964. Chinese.

28. Tewarson, A. Generation of heat and chemical compounds in fires. In *SPFE Handbook of Fire Protection Engineering*, Society of Fire Protection Engineers, Boston, 1988.
29. Trunkey, D.D. Inhalation Injury. *Surg. Clin. North Am.*, 58:1133-1140, 1978.
30. Zheda, J.K. Constructional changes of inner surface of air passage during aspiration of hot steam. *Anesthesiology*, 26: 1070-1074, 1977. Japanese.
31. Zhen-rong, G., Zhi-yong, S., Oppenheimer, L., Hoppensack, M., & Schneider, U. Pulmonary oedema in isolated lung lobe after inhalation injury. *Burns*, 17(6): 468-472, 1991.