DRUG RECOGNITION EXPERT (DRE) VALIDATION STUDY

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ABSTRACT

The method, procedures, and findings of a study of the scientific validity of an established Drug Recognition Expert (DRE) program in Arizona are reported. The DRE methodology for detecting and classifying suspected drug-impaired drivers was applied by trained officers of the Phoenix Police Department. The program was supported by comprehensive drug testing by the Arizona Department of Public Safety Crime Laboratory.

Study data were Drug Influence Evaluation records for 500 suspects who were evaluated over a 53 month period and the corresponding toxicological analyses of the suspects' specimens. The study used data base software developed for DRE data by the Southern California Research Institute.

The DREs' decisions about suspects' drug impairment status and their identifications of drug categories were highly accurate. Signs and symptoms, which were associated with specific drug categories, included dilated or constricted pupils, horizontal gaze nystagmus, and suspects' time estimates. Arrestees' characteristics and drug choices were examined. It is concluded that the DRE program, supported by the toxicology laboratory, is a valid method for detecting and classifying drug-impaired individuals.

Keywords: Drugs and Driving Toxicological Analysis Drug Recognition Expert (DRE) Program Drug Evaluation and Classification Program (DECP)

TABLE OF CONTENTS

_

ACKI	NOWLEDGEMENTS	/11
EXEC	CUTIVE SUMMARY	iii
· I.	PROBLEM STATEMENT	1
11.	HISTORY OF THE DRUG RECOGNITION EXPERT PROGRAM	1
	A. The Los Angeles Problem	
	B. The National Problem	2
	C. The DRE Program in Arizona	
		•
111.	LEGAL CHALLENGES	4
116.		T
IV.	SCIENTIFIC STUDY OF THE DRE PROGRAM	л
ΕΨ.,		+
V		c
V.	METHOD AND PROCEDURES	
	A. Study Records	
	B. Drug Recognition Experts	
	C. Drug Evaluation Procedures	
	D. Toxicological Analysis of DRE Cases	
	1. Introduction	-
	2. <u>Screening</u>	
	3. <u>Confirmation</u> 1	
	E. Data Base Entry	
	F. Data Summary and Analysis	6
VI.	FINDINGS	6
	A. Time Period and Number of Records	6
	B. Arrestee Characteristics	9
	C. DREs and Evaluations	4
	D. Toxicology Reports and DRE Opinions	6
	E. Toxicology Findings 2	
	1. <u>Positive Toxicology Specimens</u>	
	2. <u>All DIE - SER Records</u>	
	F. Signs and Symptoms and Drug Identification 4	
	1. Eve Signs	
	2. <u>Vital Signs</u>	
	3. <u>Time Estimates</u>	
	G. Arrestees' Drug Choices	
	G. Arrestees Drug Choices	ⁱ O
	DISCUSSION AND CONCLUSIONS	
VII.) [
OFFERENCIA	F0	
	ES	
	ES	8
	Roster of DREs	
· II.	DRE Court Cases and Hearings	
111.	DRE, Laboratory, and Data Base Forms	
IV.	Directory of Data Base Records	
V.	"Other" Reported Drugs	

iv

TABLE OF TABLES

	<u>.</u>	Page
1A	Radioimmunoassays	. 11
18	Index of Routine GC-MS Confirmatory Procedures	. 14
1C	Current Blood GC-MS Confirmatory Procedures	. 15
2	Age, Gender and Ethnic Distributions	. 21
з	Positive Toxicology: Ranks for Nine Drugs	. 30
4	Number of Drugs Detected, by Gender and Ethnic Groups	. 31
5	DRE Identifications of Drug(s), by Specimen	. 34
6	DRE Identification of Drugs, by Number of Drug Categories per Specimen	. 35
7	DRE Correct Identifications and Misses, by Drug for 668 Drug Detections in 416 Specimens	. 41
8	Eye Signs Observed during Drug Influence Evaluations	, 47
9	Mean Blood Pressure and Pulse Rates as Measured during Drug Influence Evaluations	. 49

.

. .

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-

- . . .

TABLE OF FIGURES

	•	_
		<u>Page</u>
1	SCRI Study Activities	7
2	DRE Evaluations by Month	. 17
3	DRE Evaluations by Year	. 18
4	Evaluations Conducted by 37 DREs	. 20
5	Ages, 500 DUID Suspects	. 22
6	500 Arrestees, Ethnic Groups	, 23
7	Drugs Detected in Specimens	. 29
8	Drug Identification, by Specimen	. 36
9	Percent Correct Identifications and Misses by Drug Category	. 37
10	DRE Identification of Drugs, by Drug Category	38
11	DRE Identification of Drugs (Multiple Drugs per Specimen)	. 42
12	DRE Measurements of Pupil Size, Single Drug Specimens	45
13	DRE Measurements of Pupil Size, Multiple Drug Specimens	46
14	Distribution of Positive BACs	. 54

.

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The DRE program requires the cooperative efforts of many individuals in law enforcement, the laboratory, and state and federal agencies. DRE-trained police officers are the foundation of the program, and we congratulate all of them for their achievements. In particular, we thank the Phoenix Police Department for agreeing to be the subject of study and for providing copies of Drug Influence Evaluations. Without the extensive contributions of Chief Dennis A. Garrett, Sergeant Dick Yost, Phoenix DRE Coordinator, Lt. Joe Klima, past Phoenix DRE Coordinator, and Officer Gary Huebner, the study could not have been accomplished.

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DPS Director Colonel F.J. "Rick" Ayars; DPS Assistant Director Lt. Colonel G.W. Ross; Arizona DRE Coordinator Vern Alley; and DPS Laboratory Supervisors Clifford C. Webber, Clifton Vander Ark, and Robert A. Jarzen.

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DRUG RECOGNITION EXPERT (DRE) VALIDATION STUDY

EXECUTIVE SUMMARY

In a research project sponsored by the Arizona Governor's Office of Highway Safety and supported by the Arizona Department of Public Safety (AZ-DPS) and the Phoenix Police Department (PPD), 500 records from an established Drug Recognition Expert (DRE) program were analyzed. Data base management and data analysis were conducted by the Southern California Research Institute (SCRI).

The study objectives were to evaluate the validity of the DRE methodology with records from an established program, to examine relationships between drug signs and symptoms and drug presence in specimens, and to study arrestee characteristics and drug choices.

<u>Section One</u>, the Problem Statement, describes the law enforcement problem which led to the development of a DRE program. An arrestee's low or negative breath alcohol test indicates that observed impairment is not due to alcohol. The officer must then make a decision whether to arrest or release, given that the impairment has some other cause. At issue is whether the decision will be made by an officer who has no specialized knowledge of drug effects or an officer who has been trained to recognize drug signs and symptoms.

<u>Section Two</u> briefly traces the development of the DRE program from its origin in Los Angeles to its application in Arizona and other states. The training program's initial development was within the Los Angeles Police Department (LAPD) with the assistance of scientists, physicians, and other experts. It evolved into a rigorous course of instruction in which officers are trained to recognize behaviors and physiological states associated with seven categories of psychoactive drugs. They perform a systematic, standardized 12-step evaluation to determine:

- whether a suspect is impaired;
- (2) if impaired, whether the impairment is related to drugs; and
- (3) if drugs, which drug category or combination of categories is present.

The program attracted widespread interest, and the National Highway Traffic Safety Administration (NHTSA) sponsored a laboratory study and a field study to examine the validity of the methods. NHTSA subsequently initiated DRE training for qualified agencies nationwide. Active units now exist in 24 states and the District of Columbia.

The DRE program was implemented in Arizona in 1987, and officers from 25 law enforcement agencies have been trained. There are 163 certified DREs statewide, with nearly 50 at both PPD and AZ-DPS.

Specimens obtained from arrestees were submitted to the AZ-DPS Central Regional Crime Laboratory for toxicological analysis. The laboratory provides scientific support for DRE units in all Arizona agencies (except the Mesa Police Department which has its own toxicology laboratory).

<u>Section Three</u> considers legal challenges to the DRE program. As expected, the validity and reliability of the methodology have been questioned. To date, the courts have supported the program.

<u>Section Four</u> discusses the specific purposes of this study. The findings provide information about:

Performance (accuracy, selectivity of DRE opinions)

A large portion of the data and analysis from this study focuses on the relationship between DRE opinions and laboratory results. Analysis of specimens provides objective corroboration of DRE opinions and the data which are necessary to assess the validity of the methodology.

Scientific validity of DRE methods

Study findings specifically address the question, "Do the DRE methods accomplish their stated purpose, i.e., the correct identification of drug impairment, as demonstrated by DRE opinions and specimen analyses?"

Types of drugs used by drug-impaired suspects

Information about drugs, drug combinations, and drug concentrations in specimens, which accumulate and change over the life of the DRE program, assists police agencies and laboratories to allocate resources effectively.

Signs and symptoms vs drug presence

A drug recognition methodology must be based on observable signs and symptoms which are demonstrably valid. A key focus of this study, therefore, has been the examination of evaluation data in relation to the specific drugs reported from specimen analysis. Note also that the DRE evaluations provide an otherwise unavailable means to study drug effects over a wide range of dose levels and drug combinations.

Socioeconomic factors

Drug availability and cost, weather, seasonal, entertainment, and athletic events, and the general economy are just some of the variables which may exert significant influence on drug use behaviors, which in turn affect DRE activities. A unit's activity also reflects agency policies and personnel, as well as the maturity of the program. Awareness of the influence of these variables is important for effective program management.

Program benefits vs costs

A DRE program's primary objective is to facilitate the enforcement of traffic safety laws, thereby reducing injuries, fatalities, and property damage. In the studied program, at least 378 drivers were removed from the roadway and prevented from driving in an impaired state. The safety benefit of DRE, however, is not without cost. The program makes significant demands on the police agency, and generates a requirement for specimen analysis which may tax laboratory resources. Costs may prove to be a formidable challenge to the DRE program.

Section Five describes the study method and procedures. A grant of funds was awarded in April 1993 by the Arizona Governor's Office of Highway Safety. The DRE records of PPD and the corresponding AZ-DPS toxicology reports were retrieved, copied and forwarded to SCRI. The 500 records represent the entire work product of the PPD DRE unit, and the sample contains no known bias. The cases meet the following criteria: 1) A driving-under-the-influence (DUI) suspect was evaluated; 2) the evaluation was performed by a certified DRE; and 3) the specimen obtained from the suspect was analyzed by the AZ-DPS Central Regional Crime Laboratory.

The DREs performed the 12-step evaluation in accordance with the program's national standards. The laboratory screened specimens by a comprehensive drug testing protocol and confirmed positives for forensically important substances by gas chromatography-mass spectrometry.

Data were entered into a computer data base, using software specifically developed for DRE records by SCRI under funding from the National Institute on Drug Abuse. Printed summaries of data for each arrestee were generated and checked for accuracy against source documents. Data summaries were obtained with the data base count capability, and analyses proceeded via logical interrogations of the data base and calculation of appropriate statistics. The data base resides in a computer dedicated to Arizona data.

<u>Section Six</u> reports study findings. On average, 9.4 evaluations were performed each month during the 53 month period of the records. There were more than three times as many male as female arrestees. In terms of 1990 census data for Phoenix, Hispanics are underrepresented and Caucasians are overrepresented. The distributions of licensed drivers or registered car owners would be more relevant comparison data but are not available.

Four drug categories appeared most often in specimens: depressants, narcotic analgesics, marijuana, and stimulants. Thirty DREs had examined suspects who had used drugs in one or more of these categories. Eighteen officers had encountered four categories, and seven officers had encountered five. DREs evaluate suspects who are under the influence of PCP, hallucinogens, or inhalants less frequently, but because of the obvious and unique signs and symptoms of these drugs, loss of proficiency in identifying them is not expected to be a problem.

DREs recognize seven drug categories, but the specimen analysis identifies specific drugs and metabolites. This difference is a key to understanding study findings. The laboratory reported 813 drugs in the 500 cases. There was one drug in 163 specimens, two or more drugs in 253 specimens, and no drug in 68 specimens. Sixteen arrestees refused to provide a specimen.

Of the 416 specimens for which the laboratory reported one or more drugs, the DREs correctly identified at least one drug in 378 specimens (91%). The laboratory identified at least one drug in support of the DRE opinion in 83.5% of cases for which the DREs identified one or more drug categories. Drugs were not found in specimens obtained from 26 individuals who were judged by the DREs not to be under the influence of drugs.

Preliminary investigation showed selected signs and symptoms to be uniquely related to the presence of specific drugs. The effects of narcotic analgesics and stimulants on pupil size were marked, confirming that pupil size is a reliable indicator for those categories. Horizontal gaze nystagmus was associated with benzodiazepines, barbiturates, and phencyclidine. Suspects' time estimates were related to type of drug, and drug effects on pulse and blood pressure were discernible as mild but real changes.

In order of decreasing frequency, marijuana, cocaine, benzodiazepines, morphine, methamphetamine, codeine, barbiturates, and phencyclidine were found in specimens. Illegal drugs predominated, but prescription drugs (benzodiazepines, butalbital, carisoprodol, and several narcotic analgesics) were also important. Cannabis emerged as the leading drug among men, benzodiazepines as the leading category among women. Impairment attributable solely to antihistamines or tricyclic antidepressants was infrequent.

<u>Section Seven</u> offers conclusions and interpretations of study findings. DRE opinions identified and classified drug-impaired drivers with a high level of accuracy. DRE positive opinions, which were entirely unsupported by analysis of a specimen, were few in number.

In terms of safety objectives, it should be noted that most of the 500 drivers could not have been arrested without the evidence of impairment obtained from the DRE evaluation, as corroborated by laboratory analysis of a specimen. Slightly less than one third of the arrestees had consumed alcohol, and only 5% had BrACs of 0.10% or higher.

The major conclusions of this study are:

- The DRE program is a valid method for identifying and classifying drugimpaired drivers.
- Certified DREs recognize drug-impairment and identify the category of drug(s).
- Observable signs and symptoms are associated with specific drugs.
- Monitoring DRE opinions and laboratory results will facilitate program management.
- The DRE program requires scientifically sound support by the laboratory.

I. PROBLEM STATEMENT

The ease of obtaining breath specimens together with the immediacy and low cost of breath alcohol concentration (BrAC) analysis have made it possible to estimate the prevalence of alcohol use among driver populations. As a consequence, the contribution of the single substance, alcohol, to traffic injuries and fatalities is reasonably well understood. Much less is known, or is likely to be known by the same methods, about other potentially impairing drugs.

The analysis of urine specimens can determine that a drug or metabolite is present, providing evidence that some unknown amount of drug was used at some unspecified time in the relatively recent past. This information alone, however, does not support estimates of drug prevalence in driver populations; i.e., it does not demonstrate conclusively that potentially impairing drugs were active in the driver at the time of driving. Such estimates require blood specimens, which are difficult to obtain and costly to analyze. Thus, data concerning the number of driving is sparse. Furthermore, the relationship of blood drug concentrations and impaired driving skills has not been established for many potentially impairing substances. Efforts to determine the role of drugs in traffic crashes continue, using a number of different methods (1, 2).

With or without information about the number of offenders or the causes of impairment, traffic officers are required as a routine duty to detect, test, and arrest impaired drivers. Notwithstanding the lack of scientific data, validated procedures, or department policy, officers are obliged to make timely decisions on a daily basis. In the case of alcohol, the suspect may or may not display gross signs of impairment, but breath test results provide immediate support for the decision to arrest or release. In contrast, if a zero or low BrAC suggests that other drugs may be impairing the driver, there are no immediate chemical test results to support a decision. An arrest/release decision must and will be made; the only question is whether it will be made by a traffic officer who has no specialized knowledge of drug effects or whether it will be made by an officer who has been trained to recognize the signs and symptoms of drug impairment.

II. HISTORY OF THE DRUG RECOGNITION EXPERT PROGRAM

A. The Los Angeles Problem

During the 1970's, Los Angeles Police Department (LAPD) traffic officers encountered an increasing number of obviously-impaired drivers whose BrACs were zero or low. The problems in evaluating, arresting, and prosecuting such drivers were the impetus for the development of a Drug Recognition Expert (DRE) methodology. A training program originated within the department, and with the assistance of scientists, physicians, and other experts, it evolved over a period of several years into a rigorous course of instruction. It is designed to train officers to recognize behaviors and physiological states associated with seven categories of psychoactive drugs.

DRE-trained officers developed the knowledge and skill which enabled them to accurately identify drug-impaired drivers, as corroborated by laboratory analysis of urine or blood specimens. Los Angeles courts began to accept their expert testimony, the number of filings of drug cases increased, the number of guilty pleas increased, and the amount of time officers were required to be present in court decreased.

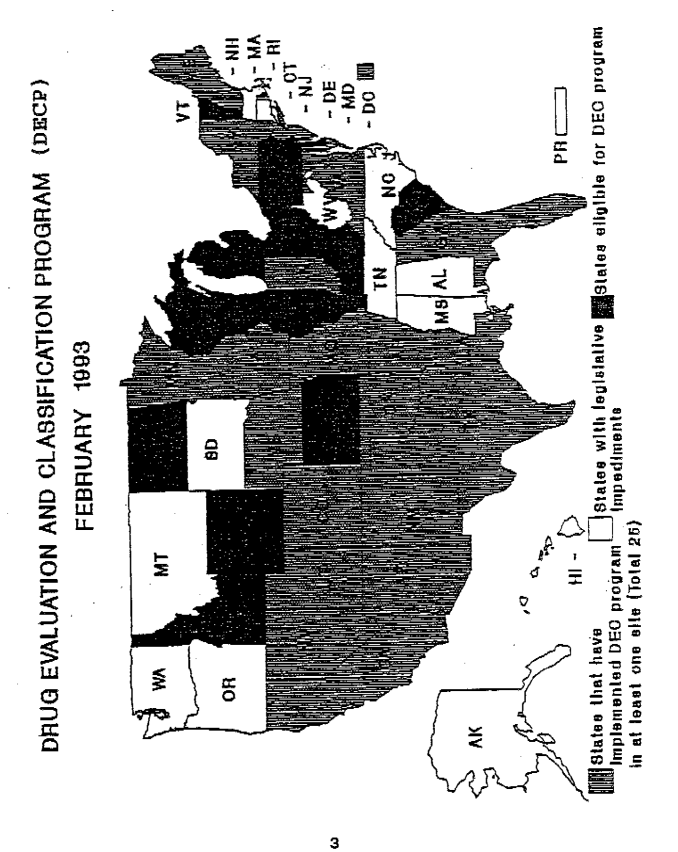
B. The National Problem

Drug use was not a problem which existed only in Los Angeles, nor was the need to properly identify, arrest, and charge drug-impaired arrestees unique to LAPD. Not surprisingly, the apparent success of the DRE program attracted widespread interest. In response to that interest, the National Highway Traffic Safety Administration (NHTSA) and the National Institute on Drug Abuse sponsored a study at Johns Hopkins University (3) to examine the validity of the methods. In a laboratory experiment, 80 subjects who had been administered a drug (amphetamine, marijuana, diazepam, or secobarbital) were examined by four LAPD DREs, using a standardized, abbreviated examination. The DRE identifications of drugs were correct for 80%, 97.5%, and 92.7% of subjects dosed with stimulants, marijuana, and depressants, respectively.

Similarly, in a 1985 field study, 25 LAPD DREs were highly accurate with regard to suspected drug-impaired drivers in the City of Los Angeles (4, 5). DREs correctly identified at least one drug in 87% of their evaluations and were correct in 94% of the cases where they judged a driver to be impaired by a drug other than alcohol.

NHTSA subsequently undertook a program to make DRE training available for qualified agencies throughout the United States. In cooperation with LAPD, they further developed the training curriculum, including instructor and student manuals, and other teaching materials. Initial DRE units were established in Arizona, Colorado, New York, and Virginia.

With overview by a Technical Advisory Panel and administration through the International Association of Chiefs of Police, the program continues to evolve. As can be seen in the figure which follows this page, active units of what is now called the Drug Evaluation and Classification Program (DECP) have been established in 24 states, the District of Columbia, Australia, Norway, and Canada. Approximately 3000 DREs and 800 instructors have been certified (6).



C. The DRE Program in Arizona

The training of Arizona DREs began in Los Angeles in 1987. Fourteen officers were trained during that year, as were two prosecutors and two scientists from the Arizona Department of Public Safety (AZ-DPS) Crime Laboratory. The training of officers, prosecutors, and crime lab personnel continued in Los Angeles into 1988. Beginning in 1989 and continuing in 1994, one (sometimes two) DRE schools have been conducted each year in Arizona.

A few Arizona candidates who attended a DRE school did not achieve certification, and a few DREs have lost their certification status. De-certification typically has occurred because an officer became inactive as a DRE as a result of transfer or promotion. At the present time, 163 law enforcement officers statewide are certified DREs. The Phoenix Police Department (PPD) currently has 47 DREs, including four supervisors.

The AZ-DPS Crime Laboratory provides toxicology support to all DRE agencies except Mesa Police Department, which has its own crime laboratory. The AZ-DPS Laboratory was established in 1969 and became a full service laboratory system with regional laboratories in Phoenix, Tucson, Flagstaff, and Mesa. Toxicological analysis of drugs is performed at the Central Regional Laboratory in Phoenix which serves over 250 city, county, state, federal, and tribal agencies in the state.

III. LEGAL CHALLENGES

As expected, defense attorneys in a number of jurisdictions have challenged the validity and reliability of the DRE methodology. Typically, they have moved to suppress evidence from DRE evaluations under the Frye standard. A list of DRE hearings and cases appears in Appendix II. To date, the courts have supported the program, but additional legal challenges are expected.

IV. SCIENTIFIC STUDY OF THE DRE PROGRAM

Socioeconomic variables exert significant but often unrecognized and unmeasured influence on drug use behaviors, which then affect the activities of a DRE unit. The drug evaluations conducted by DREs reflect the number of officers assigned to traffic duty and the number of drug-impaired drivers on the roadway. The latter is related to many variables, including drug availability and cost, season and weather, entertainment and athletic events, and the general economy. Also, a DRE unit's activity inevitably is a function of agency and laboratory policies, as well as the unit's personnel at a specific time. A new program has different performance characteristics than a mature program, but whether the changes which occur over time will be a net gain or loss is not always predictable. To some extent, conditions will be unique to the site. For example, a diminution (if any) of the enthusiasm which characterizes new programs can reasonably be expected to be offset by gains in skill and experience. Whether benefits actually do accrue, however, depends on a number of local variables, including whether the program continues to be supported within the agency, by the laboratory, by prosecutors, and by the courts.

A retrospective study examined the performance of Arizona DREs, initially with 185 cases with subsequent expansion to 341 cases (7, 8). An 86% rate of correct identifications (drug subsequently found in a sample of the suspect's urine) is remarkably close to the overall correct detections in the Los Angeles field study (4, 5). A study of 526 Arizona cases also has been reported (9). Data from DRE programs in California, Texas, and Minnesota demonstrate similar rates at 88.2%, 81.3%, and 84.5%, respectively (10, 11, 12).

The DRE program is designed to identify suspected drug-impaired drivers, thereby making it possible to remove them from the roadway. A program benefits the agency and the community, not only in traffic safety but in drug traffic and crime suppression as well. These are worthy objectives, but they are not without cost. A DRE unit places high demands on a department initially for officer training time and subsequently for duty time. Frequently, laboratories are taxed as they stretch resources to handle the additional urine and blood specimens that the program generates. Within a difficult economy and a climate of accountability, non-productive DRE units and inefficient laboratories likely will come under close scrutiny. Cost may prove to be the most formidable challenge to the DRE program.

In addition to providing data to answer questions about costs vs benefits, evaluation of DRE units will facilitate effective program management. The data will enable program coordinators to examine differences in units' activities as a function of time, location, staffing, and other variables. It will provide useful feedback on performance to the DREs themselves, and will serve as a source of scientifically sound data for the purpose of responding to legal challenges.

There is yet another reason why the records merit study. The body of drug information, which law enforcement needs, is woefully incomplete. The scientific literature about drug effects on performance and drug signs and symptoms is and likely will continue to be limited. Unlike the single substance, alcohol, there are many drugs, and the research community is unable to examine all potentially impairing substances, all dose levels, and all drug-drug, drug-alcohol combinations. Furthermore, scientific study frequently is not designed to obtain and/or report the specific data needed by law enforcement. Research which requires the administration of dangerous substances to human subjects is restricted by ethical, safety, and legal constraints. Arrestees, in contrast, are not constrained by anything other than drug availability and their own choices. They sometimes are found to have ingested illicit and/or therapeutic drugs in dangerously high amounts and in unusual combinations. In such cases, the DRE gathers data which are not available elsewhere. The records, presently residing in the files of DRE units nationwide, are an underutilized resource.

To facilitate access to the information contained in Drug Influence Evaluation (DIE) records, data base software (NIDABASE) was developed by the Southern California Research Institute (SCRI) under funding from the National Institute on Drug Abuse (13). The study described in this report used that software to examine Arizona DIE records:

- 1) for scientific purposes;
- 2) to provide data relevant to legal issues;
- to provide information about DRE performance to state and local coordinators and to the DREs;
- to examine the relationship of signs and symptoms and the presence of a drug or drugs in urine; and
- 5) to establish an evaluation mechanism in the interest of program accountability.

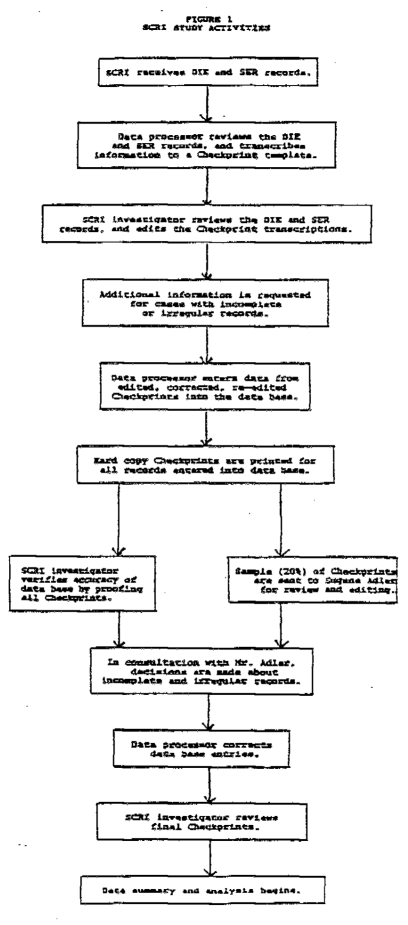
V. METHOD AND PROCEDURES

Study activities are graphed in Figure 1. A grant of funds from the Arizona Governor's Office of Highway Safety was awarded in April 1993. Records were received by SCRI in August 1993 at which time study activities were initiated at that site. Data analysis was completed in March 1994. This document reports study findings and completes the activities of this phase of study.

A. Study Records

Study data were obtained from Drug Influence Evaluation (DIE) records and the associated DPS Scientific Examination Reports (SERs) for suspects examined during the period January 1989 through May 1993. The total work product of the Phoenix Police Department DRE program over a 53 month period was retrieved, and the sample contains no known bias. The cases meet the following criteria:

- A DRE evaluated a driving-under-the-influence (DUI) suspect;
- The evaluation was performed by a certified DRE. (Evaluations performed by certification candidates during training were excluded.); and
- A specimen obtained from the suspect was analyzed by the AZ-DPS Central Regional Laboratory.



B. Drug Recognition Experts

The evaluation forms, which can be seen in Appendix III, are the records of examinations of suspected drug-impaired drivers by certified DREs. Taking the latter part of the study (1992-93) as the point of reference, the officers who conducted the evaluations had served with the department ten years and had three years' DRE experience, on average.

C. Drug Evaluation Procedures

DRE examinations typically are requested by an arresting officer after he/she has obtained a breath test result which proves to be inconsistent with the observed driving and behavioral impairment. The examinations require as much as one hour's time, and are conducted most frequently in station houses where suspects are transported by the arresting officer. If the DRE is also the arresting officer, some preliminary information is obtained at roadside. When accident-involved suspects are transported to a hospital, a partial evaluation is conducted at that location.

The drug evaluation is a systematic and standardized procedure, which includes the following twelve steps (14):

- 1. Breath alcohol test *
- 2. Interview of arresting officer
- 3. Preliminary examination and first pulse
- 4. Eye examinations
- 5. Divided attention tests
- 6. Blood pressure, temperature, and second pulse
- 7. Dark room examinations and ingestion examination
- 8. Examination for muscle rigidity
- 9. Inspection for injection sites and third pulse
- 10. Interrogation, suspect statements, and other observations
- 11. Integration of all information as basis for evaluator's opinion
- 12. Toxicological examination

In all circumstances, the objectives of the evaluation are to enable the DRE to determine:

- whether the suspect is impaired;
- if impaired, whether the impairment is related to drugs; and
- if drugs, which drug category or combination of categories is present.

^{*} PPD obtains breath specimens for BrAC measurement with a gas chromatograph (Intoximeter, GCI Mark IV). The instruments were maintained by the City of Phoenix Police Crime Laboratory. They were operated in accordance with AZ-DHS regulations by officers who are DHS licensed GCI operators.

D. Toxicological Analysis of DRE Cases

1. Introduction

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Study of the DRE program requires definition of the data to be examined, i.e., the Drug Influence Evaluations and the toxicology reports. A very large data set from a number of DRE sites and laboratories would provide the statistical power to examine numerous potentially important variables. It might also introduce error from significant but unrecognized differences between protocols and procedures. Mean values calculated from such heterogeneous data are potentially useful for monitoring driving-under-the-influence of drug (DUID) trends, but they do not serve an evaluation of DRE performance or the examination of the relationship of signs and symptoms with drug concentration in a specimen. To facilitate the objectives of this study, homogeneous data from a single program served by a single laboratory during a defined time period have been examined.

Numerous substances qualify as drugs of abuse, but few are actually common in DUID cases. Three illegal drugs predominated in this study: marijuana, cocaine, and methamphetamine. Knowledge has accumulated over the life of the DRE program about the specific drugs which are likely to be found most frequently in specimens obtained from DUID suspects. That knowledge aids in the appropriate utilization of laboratory resources.

Still, toxicologists confront numerous difficult decisions about specimen choices and analytical methods and schemes, as well as their ultimate philosophy of DUID case investigation. Which drugs should be tested for? Which cutoffs are appropriate? Should the screening panel be the same for all cases? Which screening positives should be confirmed, given a particular DRE opinion? When should quantitative analysis be performed?

It is imperative to find reasonable and effective answers to these questions in order to integrate toxicological support with the DRE program in a manner which significantly advances the overall goal of detecting drug-impaired drivers. The program, although systematic and standardized for the law enforcement officer, came to the toxicology laboratory somewhat like a kit requiring assembly. Both the program and scientific support continue to evolve.

Specimen choice is the subject of regular, sometimes acrimonious discussion among toxicologists. In DUID cases, the choice is constrained by legal, logistical, and budgetary issues, as well as by toxicological considerations. The quicksand of the subject matter is not germane to this report except for a brief comment on specimen choice as it applies to the study data.

Neither blood nor urine is perfect for analysis. Each has advantages and disadvantages, but the AZ-DPS Laboratory's recommendation to all its user agencies is that urine is the preferred sample to be routinely obtained. Urine can be comprehensively analyzed at reasonable cost for most substances involved in DUID cases. Toluene is an exception, and blood specimens are recommended when inhalants are suspected.

The AZ-DPS Laboratory acknowledges the occasional need for quantified drug and metabolite concentrations in blood. In serious accidents with injuries and fatalities, particularly if a driver's injuries limit the opportunity to directly observe drug signs and symptoms, the collection and analysis of both blood and urine may be recommended. Routine analysis of both, however, is typically not an option, and a choice must be made between the two fluids.

The forensic analysis of drugs in urine or blood must be as comprehensive, accurate, and systematic as possible. The design of the DPS Laboratory's toxicological protocol meets these criteria and permits scientifically valid evaluation of the DRE program. During the 53 month period from which the study data came, no significant changes were made in DRE evaluations, and only minor changes and improvements (as noted) were made in the toxicology protocol.

Strong quality assurance and reliable performance are prerequisites for providing accurate, qualitative toxicological data for both the support and the evaluation of a DRE program. The AZ-DPS Laboratory's quality assurance program, which predates DRE, incorporates quality control into all analyses. The lab also maintains a proficiency testing program (external and in-house), and it performs continual casework review to assure quality. External evaluation of lab performance is necessary. Note that the Arizona DPS Laboratory was accredited by the American Society of Crime Laboratory Directors (ASCLD) in 1982 and has maintained its accreditation status since that date.

From a broader view of laboratory assessment, the following professional organizations and agencies serve as references and standard bearers for laboratories involved in the DRE program nationwide: ASCLD, American Academy of Forensic Sciences, Society of Forensic Toxicologists, National Institute on Drug Abuse, and the College of American Pathologists. Also, the Toxicologists Advisory Group of the Drug Evaluation and Classification Program, which meets periodically with NHTSA, has produced a site assessment protocol for the evaluation of laboratories seeking entrance into the DRE program.

2. <u>Screening</u>

The increased volume of DUID cases generated by trained officers is compatible with the trend toward automation in the laboratory. DRE cases are particularly amenable to systematic, automated screening. The screening analysis must be as comprehensive as possible with few significant analytical blind spots. The objective is to achieve a high detection rate without allocation of laboratory resources to rare or forensically unimportant substances. Secondary screening by gas chromatography with flame ionization detectors (GC-FID) was performed throughout the entire study period (15). The rules governing secondary screening were as follows:

- a. IF a DRE opinion includes depressants (other than alcohol) AND the RIA screening for barbiturates and benzodiazepines is negative (or does not lead to a confirmed depressant), THEN secondary screening for other depressants shall be performed.
- b. IF a DRE opinion includes narcotic analgesics AND the RIA screening for opiates is negative (or does not lead to a confirmed opiate), THEN secondary screening for other narcotic analgesics shall be performed.
- c. IF analysis of a miscellaneous drug (such as carisoprodol, ethchlorvynol, or meperidine) is specifically requested or indicated by the case history, appropriate screening for that substance shall be included in the case analysis.

3. Confirmation

The detection by screening of significant or potentially significant drugs was followed with confirmation by appropriate gas chromatography-mass spectrometry (GC-MS) procedures. The confirmation of so many substances in the numerous specimens generated by a mature DRE program is a formidable task, and it requires a set of confirmatory procedures designed to achieve the best compromise between sensitivity, simplicity, and efficiency.

Sensitivity entails sophisticated techniques, as does automation, but the application of a limited set of routine procedures can facilitate efficiency. Toward that objective, the number and complexity of confirmatory GC-MS procedures were minimized, and the analytical scheme was made as simple as possible. The GC-MS procedures for urine, which had been established prior to the period of this study, were not altered except for improvements in the sensitivity of the opiate and benzodiazepines procedures.

The simplest procedure was a rapid liquid-liquid basic extraction followed by full scan GC-MS in the electron ionization (El) mode. Although almost any conventional basic extraction can work, convenient "TOXI-A" extraction tubes and "TOXI-A" discs (ANSYS Inc, formerly Toxilab Inc) were employed. Some case specimens required no further confirmatory analysis. This "TOXI-A" procedure sufficed for routine confirmation of phencyclidine, carisoprodol, meprobamate, and miscellaneous bases such as tricyclic antidepressants.

The "TOXI-A" procedure was generally inadequate for the routine analysis of methamphetamine, benzoylecgonine, opiates, and benzodiazepines. In some

The primary screening process was a battery of seven radioimmunoassays (RIA), DPC Corporation, routinely applied to all incoming urine specimens (Table 1A). The battery was applied regardless of requests for less extensive, specific analysis, which may have accompanied the submission of the sample. For blood, a similar routinely-applied RIA battery (excluding cannabinoids) was implemented during the study period (January 1990).

TABLE 1A

Radioimmunoassays				
Cutoff, Urine (ng/mL)	Cutoff, Blood (ng/mL)			
50 (a)	~ -			
300	50			
500 (b)	25			
150	10			
100	100			
100	50			
25 (c)	10			
	Cutoff, Urine (ng/mL) 50 (a) 300 500 (b) 150 100 100			

(a) This cutoff was reduced from 100 to 50 in 1990.

- (b) This assay is less than 5% cross reactive to the l-isomer of methamphetamine.
- (c) A sudden, unexplained decrease in phencyclidine cases occurred in 1990. Phencyclidine was eliminated from the RIA battery in January 1993, and since that time has been tested only by request.

The RIA battery does not detect all depressant and narcotic drugs, and secondary screening is sometimes required. In Arizona DUID cases, the most significant other drugs requiring secondary screening have been:

- carisoprodol and its metabolite, meprobamate
- methadone and its metabolites.
- propoxyphene and its metabolites
- meperidine
- tricyclic antidepressants (especially amitriptyline)
- antihistamines

cases, however, it did provide confirmation of methamphetamine, or free cocaine and/or <u>methyl</u>ecgonine. Overall, this is an extremely rapid, simple procedure which extracts many drugs and metabolites.

The confirmations of methamphetamine, cocaine/metabolites, opiates, and benzodiazepines were considered negative only after analysis by one of the specialized procedures discussed below with negative results. The TOXI-A procedure usually confirmed barbiturates, but attempts to confirm barbiturate positives were not considered exhausted until a special acidic extraction (employing "TOXI-B" tubes) was performed.

Analysis of benzodiazepines and opiates required hydrolysis, derivatization, and the selected ion monitoring (SIM) mode. If desired, the analysis of both opiates and benzodiazepines could be batched, sharing the same extraction and derivatization after providing each analysis with the appropriate internal standards, blanks and controls. The GC-MS Data System was programmed to monitor various combinations of selected ions during designated time windows throughout the run. In this way, eight benzodiazepines and/or metabolites, and six opiates, were readily confirmable.

There was no difficulty in analyzing the trimethylsilyl (TMS) derivatives of lorazepam, oxazepam, temazepam, desmethyldiazepam, desalkylflurazepam, hydroxyethylflurazepam, alpha-hydroxyalprazolam, and alpha-hydroxytriazolam.

The opiates routinely analyzed as TMS derivatives were morphine, codeine, hydrocodone, dihydrocodone, oxycodone, and O-6-monoacetylmorphine (found in approximately half the cases in which morphine was confirmed).

A special extraction was necessary for THC-COOH (9-carboxy-11-nor-delta-9tetrahydrocannabinol), followed by derivatization and a reduced El scan, M/Z 200-500. Table 18 is an index of the confirmatory procedures.

TABLE 1B

Index of Routine GC-MS Confirmatory Procedures (a)

	Procedure	Int. Std.	<u>Hvdrol?</u>	<u>Deriv?</u>	<u>MS Range</u>
Α.	TOXI-A (Basics)	lprindole (b)	No	No	40-360
В.	Barbiturates	various	No	No	40-360
Ċ.	Methamphet. (c)	N-Prop. amph.	No	TFA	50-200
D.	Benzoylecg. (d)	Scopolamine	No	TMS	75-375
E.	тнс-соон	delta-8 THC-COOH	Yes	TMS	200-500
F1.	Opiates	Nalorphine	Yes	TMS	SIM
F2.	Benzodiaz.	Bromazepam	Yes	TMS	SIM

- (a) All the above procedures have in common these elements: liquid-liquid extractions; the GC column is crosslinked Phenyl Methyl Silicone 9.1 m x 0.2 mm x 0.33 mm film thickness; electron ionization mode; automated runs (autosampler), qualitative analysis; appropriate internal standards, blanks and controls.
- (b) Other internal standards, such as SKF-525, may be used.
- (c) This analysis includes ephedrine, pseudoephedrine, and amphetamine.
- (d) An alternate procedure was also used for simultaneous analysis of cocaine, benzoylecgonine, and methylecgonine.

Regarding the analysis of blood specimens submitted by DREs, radioimmunoassay, supplemented by GC-NP screening, has been effective. Blind spots for some drugs in the analytical scheme remain a concern. Solid phase or liquid-liquid extraction followed by SIM-GC-MS appears to be effective in confirming drugs of interest (Table 1C). Continuing refinement of the laboratory's procedures for blood has established effective quantitative assays, which at this time have been applied to a limited number of DRE cases.

TABLE 1C

Current Blood GC-MS Confirmatory Procedures

Procedure	Extraction	<u>Derivative</u>	MS Range
Cocaine/BE	Liq/Liq	TMS	SIM
Methamp/Amp	Liq/Liq	ŤFA	SIM
Phencyclidine	SPE (a)		SIM
Opiates	SPE	TFA	SIM
Barbs	Liq/Liq		Reduced scan
Benzodiaz.	SPE	TMS	SIM
Basics, Misc.	Liq/Liq		Reduced scan

(a) SPE (solid phase extraction) procedures were derived from Varian Corporation procedures.

E. Data Base Entry

The data base software stores pertinent DIE and SER information on a computer hard disk and prints each record as a two page summary. This study's data resides in a computer dedicated to the Arizona project. The printed summary of information for each arrestee is referred to as a checkprint (Appendix III). As can be noted by inspection of the checkprint template, arrestees' names and other uniquely identifying facts are not recorded.

The procedures for data entry and verification are graphed in Figure 1. Initially, the project data processor transcribed information contained in the DIE forms and SERs to a paper template of the checkprint. The SCRI investigator reviewed the DIE forms and SERs together with the checkprint transcription. The corrected information was entered into the data base, which assigns sequential numbers to the records.

Printouts of the checkprints were proofed by the investigator, and the data processor made needed corrections. A twenty percent sample of checkprints was drawn by taking every fifth sequential record, and copies were forwarded to Eugene Adler, DPS Laboratory, for review. Based on his review, the data processor made additional corrections to data base entries. The iterative process of proofing and correcting has produced a data base of highly accurate information.

F. Data Summary and Analysis

The Directory of Records contained in the data base appears in Appendix IV. Many of the data base entries are non-numeric (checkboxes, Yes/No, present-/absent). The data which are classificatory and nominal in character support descriptive statistics. For statistical analyses by computer, numerical data are exported from the data base to statistics programs. In addition, the program's Summary Count function is a convenient method for reporting a two-level structure of specified groups for which selected data are counted. Specified counts can be executed for all records or for a defined subset.

The Foxplus software permits direct interrogation of the data base to determine the relationships of any set of variables using commands written as logical expressions. Exhaustive exploratory analyses, which were performed using this very powerful capability, produced most of the findings reported in this document. Rank correlations and the <u>t</u> statistic have been calculated where appropriate.

VI. FINDINGS

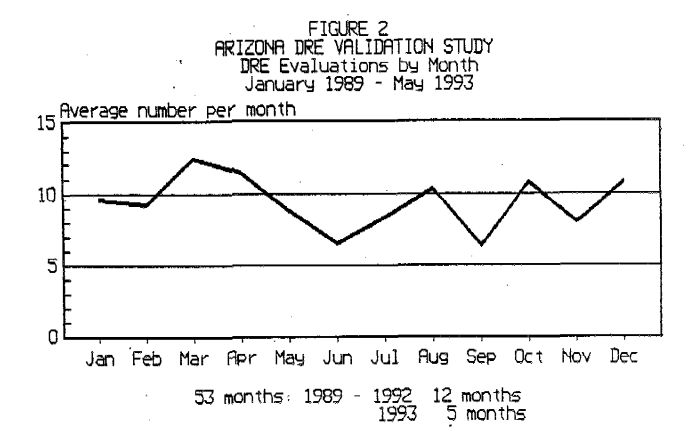
A. Time Period and Number of Records

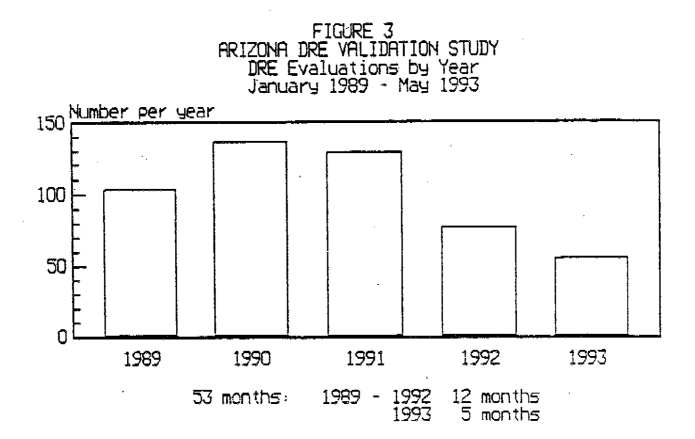
The data base covers the 53 month period, January 1989 through May 1993. It contains information obtained from the Phoenix Police Department and the Arizona DPS Laboratory with 500 DIE and SER records for 392 men and 108 women. An additional 27 records were examined but the data were not entered because the documents were incomplete.

The total numbers of records for each study year are:

1989	103
1990	136
1991	129
1992	77
1993	55

The mean number of drug evaluations performed per month across multiple years was 9.4 with a range of 6+ to 12 per month (Figure 2). In reviewing Figure 3, which graphs the number of evaluations by year, note that only 1990 and 1991 are comparable. New programs require some time period to become fully operational and 1989, the first year of full operations, may have differed from sub-





sequent years. The data base includes records for only five months of 1993, whereas records were obtained for twelve months of each of the other four years. Also, significantly fewer evaluations were performed in 1992 (1992 vs 1990 \underline{t} -3.321, p<.001; 1992 vs 1991 \underline{t} -2.575, p<.05).

During the study period, some officers were responsible for only a few evaluations whereas numerous evaluations can be credited to others. The numbers ranged from 1 to 33, with 23 DREs conducting ten or more evaluations and 14 DREs conducting fewer than ten. Among the latter were three officers who conducted one evaluation each (Figure 4).

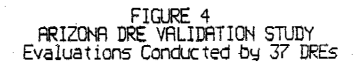
B. Arrestee Characteristics

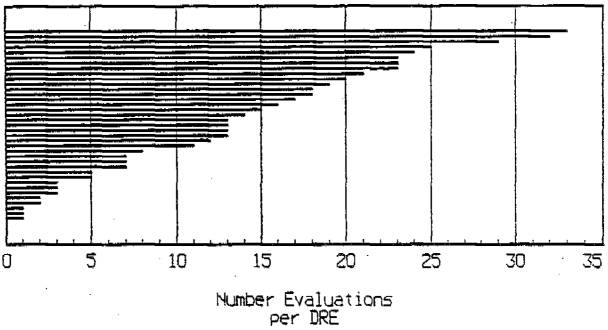
The age, gender, and ethnic characteristics of the 500 arrestees are summarized in Table 2. The arrestees were predominantly young adult males. There were more than three times as many men as women.

A wider age distribution for men than for women can be seen in Figure 5. Male arrestees were most frequently in the age group 20 - 29 years. The largest number of women were 21 - 40 years of age. Few female arrestees were under age 21, but almost 12% of the men fell in that age range. More than 5% of the men were older than age 50, and one woman was over age 60.

Almost 85% of the arrested drivers were Caucasian, 10% were Hispanic, and 6% were Black (Figure 6). No Asians were evaluated by DREs during the entire study period, nor were there any Hispanic females among the suspects. With the exception of five Black women, the female arrestees were Caucasian.

With the data at hand, it is not possible to conclude with certainty that members of one ethnic group are more or less likely than another to drive in a drug-impaired condition. If viewed in terms of the 1990 census data for the general population of Phoenix (5% Black, 20% Hispanic, 72% Caucasian), it appears that Hispanics are underrepresented and Caucasians are overrepresented in the sample of arrestees. However, the distributions of licensed drivers and/or registered car owners, data which are not available, would be more directly relevant and might or might not parallel the census data.





AGE (yrs)		All <u>stees</u>	· · · ·	<u>nales</u>	<u>N</u>	<u>lales</u>
< 21	52	10.4	7	6.5	45	11.5
21 30	190	38.0	42	38.9	149	38.0
31 - 40	156	31.2	44	40.7	112	28.6
41 - 50	71	14.2	12	11.1	59	15.1
51 - 60	14	2.8	0	0	14	3.6
61 - 70	9	1.8	1	0.9	7	1.8
Unknown	<u>8</u> 500	<u> 1.6</u> 100	_ <u>_2</u> 108	<u>1.9</u> 100	_ <u>6</u> 392	_ <u>1.5</u> 100

TABLE 2 ARIZONA DRE VALIDATION STUDY Age, Gender and Ethnic Distributions 500 Arrestees

ETHNICITY		All estees %	<u> </u>	nales	M No.	ales
Caucasian Hispanic Black Amer. Indian Not recorded	419 46 31 3 <u>1</u> 500	83.8 9.2 6.2 0.6 <u>0.2</u> 100	103 0 5 0 _0 108	95.4 4.6 	316 46 26 3 <u>1</u> 392	80.6 11.7 6.6 0.8 <u>0.3</u> 100

.

Single-drug detections are listed below:

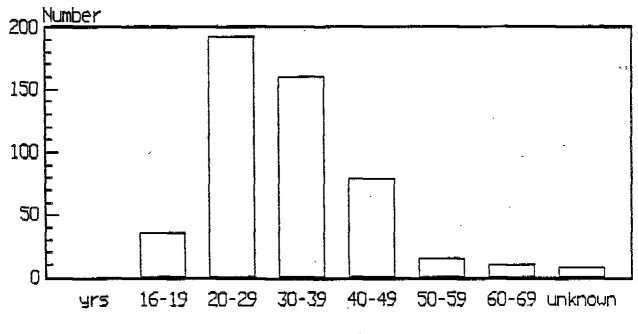
Drug	Detected Alone (no.)
Marijuana	61
Cocaine	26
Benzodiazepines	16
Methamphetamine	13
PCP	8
Barbiturates	6
Morphine	3
Codeine	1
Other drugs	<u>_29</u>
	163

In total, the detected drugs, reported in the checkprint as TOXICOLOGY RESULTS, are the following (Figure 7):

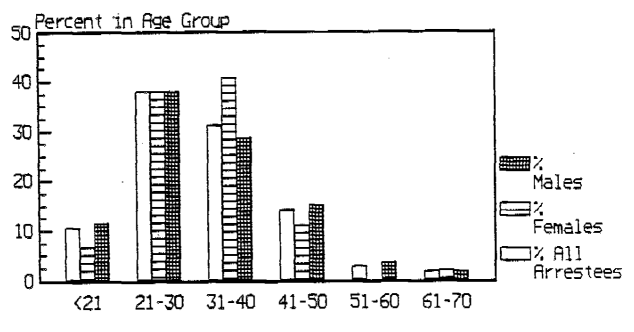
	Drug Detected (no.)
Marijuana	165
Cocaine	115
Benzodiazepines	108
Morphine	71
Methamphetamine	69
Codeine	65
Barbiturates	35
PCP	22
Amphetamine	18
-	668
Other	<u>145</u>
	813

Table 3 lists rankings by frequency of detection for the total sample for men and women. They are tabled by gender and ethnicity in Table 4. Since there were many more male than female arrestees in the sample, their drug choices dominate the overall tallies. Marijuana was the drug-of-choice for Caucasian and Hispanic men whereas benzodiazepines ranked first among women. Cocaine, codeine, and marijuana were detected with approximately equal frequency in urine specimens obtained from female arrestees. Note that the women account for 22% of total group (108 of 500 arrestees), and their specimens account for 26% of detections (209 of 813 drugs). PCP was found twenty times in urine obtained from men, but only twice in specimens obtained from women.

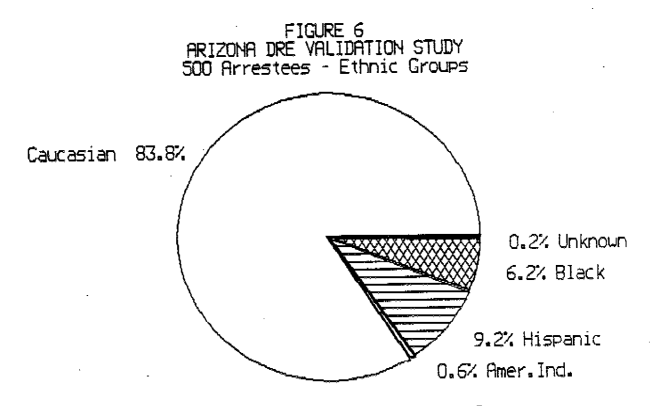
FIGURE 5 ARIZONA DRE VALIDATION STUDY Ages, 500 DVID Suspects



years



Age Groups (years)



Percent of Total Sample by Ethnic Group

With few exceptions, DREs did not record "employment status" of arrestees during the period 1989 - 1990. Although they began in 1991 to note the arrestees' occupations more frequently, the information is available overall for less than 20% of the group. With the occupation of 411 arrestees unknown, the value of the following information is extremely limited, and certainly cannot be generalized beyond the 89 arrestees to whom it applies.

	NUMBER	PERCENT
Unemployed	30	33.7
Unskilled	7	7.9
Semiskilled	18	20.2
Skilled	25	28,1
Professional	4	4.5
Student	5	5.6
Total	89	

C. DREs and Evaluations

Significant resources have been required to train Arizona officers in the DRE methodology, and it is reasonable to inquire about the benefits for law enforcement and the community at large. Is the unit meeting the objectives which underlie the adoption of DRE in Phoenix? Is the unit having an impact on traffic safety in Phoenix?

The number of DUID suspects evaluated by the unit and by individual officers can be taken as relevant measures of DRE activity. In general, arrests parallel evaluations except that evaluated drivers are not arrested if they are found to be "not impaired." Although an evaluation is requested only when there is evidence of impairment, the DRE may conclude at the end of an examination that the suspect is experiencing a medical problem, extreme fatigue, or emotional distress, and that no impairing substance is present.

When an evaluation does culminate in an arrest, the driver is prevented from crashing on that occasion. In that sense, the number of arrests is an index of the program's short term contribution to roadway safety. A more difficult query concerns the program's long term safety benefits. A satisfactory answer to that question will require analysis of a broader data set, which includes injury and fatality statistics over a longer time period.

The number of DREs who conduct evaluations over an extended period postcertification is an index of program activity. The PPD data show significant between-DRE variability. It should be kept in mind that whether a DRE does or does not examine drug-impaired drivers is related not only to the individual officer's assignments and motivation, but also to department priorities and budgets, the DRE unit policies, drug availability, drug cost, the weather, the economy, and other diverse, sometimes unrecognized influences. Such variables alter the number of drug-impaired drivers on the roadway at any given time, the number of traffic officers on patrol to detect them, and the number of DREs available to examine them. It is not possible to retrospectively identify and analyze all of these variables with available data and resources, but their impact should not be minimized.

The number of evaluations is, at least in part, a function of elapsed time since an officer's certification. As expected, an examination of the Phoenix data indicates that for most but not all officers, the premise of a time-number relationship is valid. Using the dates of first and most recent evaluations to approximate time-since-certification, it was found that the officer who conducted evaluations over the longest period of time (51 months) is also the officer with the largest number of evaluations (33). More broadly, if the analysis is restricted to those DREs who conducted ten or more evaluations during the study period, number is significantly related to time (Spearman Rank correlation, 0.67, p < .005).

Activity level is also important in terms of officers being able to maintain proficiency with DRE skills. It is an issue not only of the total numbers but of the particular drugs and drug combinations which are encountered. The study records were examined to determine how many times each DRE examined suspects under the influence of drugs in each of the seven categories. If most suspects in a particular locale are under the influence of the same drugs (marijuana or cocaine, for example), it might be possible to conclude that the DREs are very skilled in identifying those drugs, but to be uncertain about their skills with other categories.

The four drug categories which appeared most often in specimens were depressants, narcotic analgesics, marijuana, and stimulants. Thirty of the 37 DREs had examined suspects who had used drugs in one or more of these categories (1 to 15 suspects). Eighteen officers had encountered four categories, and seven officers had encountered five. Most, if not all, DREs in this study can be expected to maintain proficiency in the four most common categories.

The signs and symptoms associated with PCP, hallucinogens, and inhalants are obvious and unique and their recognition is not expected to be difficult even for officers who encounter them infrequently. It is concluded that loss of proficiency is not currently a problem for the participating DREs; if there is any risk at all, it will be limited to officers who conduct so few evaluations that they are likely to be placed on inactive status.

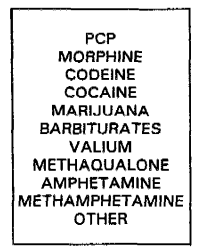
D. Toxicology Reports and DRE Opinions

An understanding of the toxicology findings, and of the DREs' opinions in relation to those findings, will be facilitated by a comparison of the DRE protocol vs the laboratory analysis. The differences between the data sources are a key to understanding the findings of this study. Reference to the checkprint template and the laboratory report in Appendix III is suggested.

A DRE identifies substances as belonging to one of seven <u>drug categories</u>. An opinion at the conclusion of the evaluation is recorded in the format illustrated below. (See page 2 of checkprint, "DRE OPINION.")

MEDICAL PROBLEM STIMULANTS PHENCYCLIDINE HALLUCINOGENS CANNABIS INHALANTS DEPRESSANTS NARCOTICS OTHER

The laboratory, however, reports the <u>specific drugs</u> which are confirmed. Positive toxicology findings are recorded in the data base in the following format. (See page 2 of checkprint, "TOXICOLOGY RESULTS.")



The important distinction is that the laboratory is able to detect and report specific drugs whereas a DRE identifies and reports substances by category. Drug signs and symptoms do not permit him/her to distinguish between morphine and codeine, for example. Based on observations only, there is no unique sign or symptom which identifies a drug as amphetamine instead of methamphetamine. In these cases, a DRE identifies and reports "narcotic analgesic" and "stimulant."

Because it is not feasible to predict trends in users' choices or to provide spaces in the data base for all possible drugs, the software limits the checkboxes (see preceding page) to those which were detected most frequently in the Los Angeles area at the time the software was being developed. Diazepam (Valium) was the most commonly-abused benzodiazepine at that time. Presently, however, other benzodiazepines are frequently detected in specimens, and the checkbox "Valium" has been used in this study for the broader category, benzodiazepines. Methaqualone appears in the checkboxes because it previously was an abused drug, but there is no occurrence of it in the data base records. For other drugs reported by the laboratory, the "Other" box was checked with the drug's name typed into the space below. Other drugs in this study are listed in Appendix V.

Note that the <u>drug</u> checkboxes account for only five of the seven <u>categories</u>. Inhalants and hallucinogens were not allotted a space, because many laboratories do not have the capability to analyze them and they are seldom reported. The inhalants reported for suspects arrested during the time period of this study have been recorded under "Other."

The following example illustrates a difference between what is recorded for a single case for the DRE opinion and for the associated toxicology result. Suppose a DRE concludes that a suspect is under the influence of a depressant; he records his opinion on the DIE form as "Depressant." He obtains a specimen and submits it to the laboratory for analysis. If the laboratory detects methaqualone, a barbiturate or a benzodiazepine, it will be specifically recorded in the data base as such. If another depressant is detected, it will be recorded as "Other."

E. Toxicology Findings

Findings from the laboratory analysis of the specimens obtained from arrestees can be summarized briefly as follows:

<u>Specimens (no.)</u>	
163	1 drug detected
253	2 or more substances detected
68	No drug detected
<u> 16</u>	Refusals (no specimens obtained)
500	

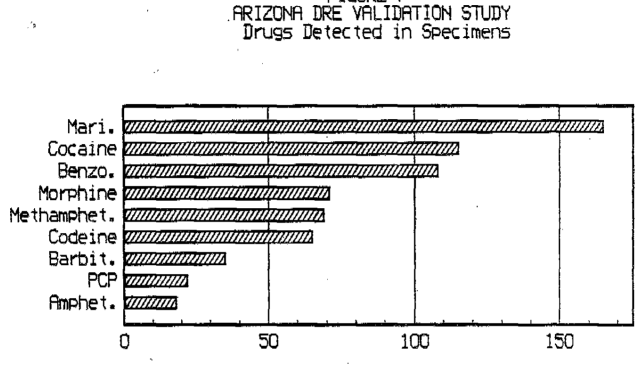


FIGURE 7

Number Times Detected

		······································			······································
	TOTAL : <u>N =</u>	SAMPLE 500_		LES <u>= 392_</u>	FEMALES <u>N = 108</u>
DRUG	<u>no.</u>	<u>Rank</u>	<u>no.</u>	<u>Rank</u>	<u>no. Rank</u>
Marijuana	165	1	144	1	21 4
Cocaine	115	2	92	2	23 2
Benzodiazepines	108	з	72	3	36 1
Morphine	71	4	55	4	16 7
Methamphetamine	69	5	52	5	17 6
Codeine	65	6	43	6	22 3
Barbiturate	35	7	17	8	18 5
PCP	22	8	20	7	2, 9
Amphetamine	<u>18</u>	9	<u>13</u>	9	<u> 5 8</u>
	668		508		160

TABLE 3ARIZONA DRE VALIDATION STUDYPositive Toxicology*: Ranks for 9 Drugs500 Arrestees

*Other drugs were identified in 145 specimens.

		ALES 108	MALES <u>N = 392</u>
	Black	Cauc.	Black Cauc. Hisp. Other
	<u>n=5</u>	<u>n=103</u>	<u>n=26 n=316 n=46 n=4</u>
DEPRESSANTS Barbiturates Benzodiaz.	0 2	18 34	0 17 0 0 1 67 4 0
NARCOTIC ANAL. Morphine Codeine	1 1	15 21	5 44 5 1 4 34 5 0
STIMULANTS Cocaine Amphetamine Methamphet	2 0 0	21 5 17	11 66 14 1 O 12 1 O 1 49 2 O
MARIJUANA	2	19	9 119 16 0
PHENCYCLIDINE	2	0	12 4 4 0
OTHER DRUGS	_2	47	<u>1 86 7 2</u>
TOTAL	12	197	44 498 58 4

TABLE 4 ARIZONA DRE VALIDATION STUDY Number of Drugs Detected, by Gender and Ethnic Group 500 Arrestees

4

The terms, which will be used to report DRE opinions as supported or not supported by analysis of specimens, are illustrated below.

And the second sec	
Hit	Drug predicted by DRE, Drug found by lab.
Miss	Drug not predicted by DRE. Drug found by lab.
False Positive (F.P.)	Drug predicted by DRE. Drug not found by lab.
Correct Rejection	No drug predicted by DRE No drug found by lab.

		τοχις	OLOGY RE	SULTS
			DRUG +	DRUG O
	2	DRUG +	нг	FALSE POS.
E I (RN	DRUG O	MISS	COR. REJECT

The DRE methodology mandates both the standardized evaluation and the analysis of a specimen. Together, the evaluation and the analysis create a balance, which is designed to identify impaired suspects (minimize misses) and, equally important, to recognize that suspects are <u>un</u>impaired (minimize false positives). False positives occur whenever:

- · the DRE misinterprets impairment signs and symptoms; or
- the DRE identifies signs and symptoms of a drug, but the limitations of the laboratory analysis result in a failure to detect it in the specimen.

Misses occur whenever:

- a suspect exhibits the signs and symptoms of a drug, but the DRE does not recognize them;
- the DRE associates a drug's signs and symptoms with another drug which is also present;
- the signs and symptoms of one drug counteract or mask the signs and symptoms of another drug; or
- the suspect was not impaired at the time of the evaluation and exhibited no signs and symptoms of impairment, but the drug or metabolite was detected in the urine specimen.

In the latter case, the DRE evaluation insures that the motorist will not be charged erroneously with being under the influence of a drug.

1. Positive Toxicology Specimens

The DRE opinions will be assessed in a variety of ways. An overview begins with 416 specimens for which the laboratory reported one or more drugs (Table 5). Looking just at those specimens which contained a drug(s), the DREs identified at least one drug in 378 specimens (91%).

2. All DIE - SER Records

In a more comprehensive analysis, DRE decisions will be assessed in terms of all data base records (Tables 5 and 6). Sixteen arrestees refused to provide specimens, and the total number of analyzed specimens for 500 suspects was 484.

The DREs identified at least one drug in 378 specimens, and drugs were not found in the specimens obtained from 26 individuals who the DREs judged <u>not</u> to be under the influence of drugs (Figure 8). Thus, the DRE decisions were supported by laboratory analysis for 404 (83.5%) of the 484 specimens, and were not supported in 80 cases (16.5%).

To more fully assess DRE performance, it is important to consider <u>how</u> decisions were right and wrong, by subsets of the arrestees, by drug category, and by other variables of interest (Figure 9). Misses or false positives occurred in 56 cases (Figure 10). Misses and false positives also occurred in combination with hits.

<u>Classification</u>	Number		
нт	184		
HIT and FALSE POSITIVE	56		
HIT and MISS	115		
HIT and FALSE POSITIVE AND MISS	23		
TOTAL with one or more HITS		378	
MISS	14		
MISS and FALSE POSITIVE	<u>24</u>		
TOTAL with no HITS		<u>_38</u>	
TOTAL: specimens in which one			
or more drugs were detected			416
FALSE POSITIVES	42		
CORRECT REJECTIONS (RULE OUTS)	<u>_26</u>		
TOTAL: specimens in which no			
drugs were detected			68
REFUSALS: no specimens obtained			16
TOTAL: arrestees			500
		•	

TABLE 5 ARIZONA DRE VALIDATION STUDY DRE Identifications of Drug(s), by Specimen *

 Classifications are per specimen with one or multiple drugs.

KEY TO CLASSIFICATIONS

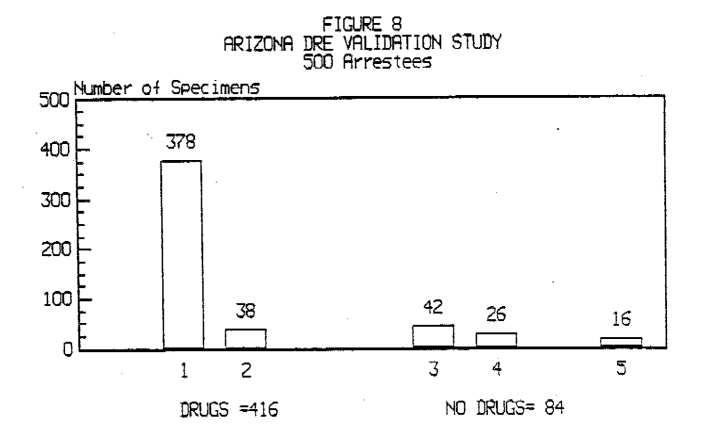
HIT MISS FALSE POSITIVE CORRECT REJECTION Drug(s) predicted and found. Drug(s) not predicted but found. Drug(s) predicted but not found. Drug(s) not predicted or found.

The DREs identified at least one drug in 378 specimens, and drugs were not found in the specimens obtained from 26 individuals who the DREs judged not to be under the influence of drugs (Figure 8). Thus, the DRE decisions were Supported by laboratory analysis for 404 (83.5%) *of* the 484 specimens, and were not supported in 80 cases (16.5%).

TABLE 6ARIZONA DRE VALIDATION STUDYDRE Identification of Drugs, by Numberof Drug Categories per Specimen

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Ţ.	1	Ħ	Hits
	2	=	Misses
!	3	<u>-</u>	False Positives
1	4	=	Correct Rejections
	5	=	Refusals

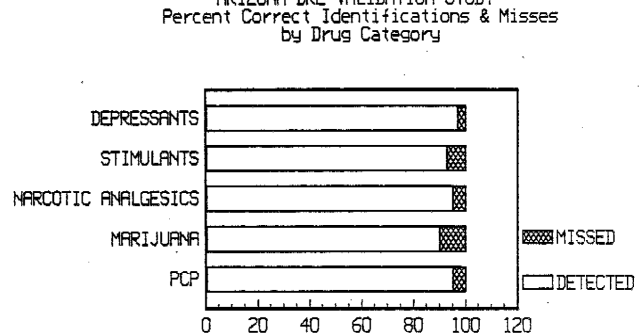
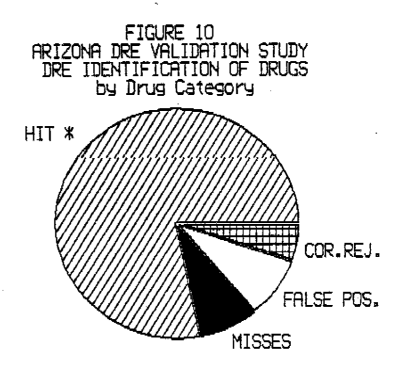


FIGURE 9 ARIZONA DRE VALIDATION STUDY Percent Correct Identifications & Misses by Drug Category

37



HIT - DRE identified at least one drug detected in analysis of specimen The laboratory detected 813 drugs (668 checkbox drugs + 145 other drugs). Table 7 displays the DRE Hits and Misses for the 668 drugs, by drug category. As can be seen, cocaine and marijuana were missed most frequently. A miss together with a hit occurred in 115 cases (Table 6). That is, the DRE identified one or more drugs but also missed one or more. In total, one or more drugs were missed in 176 decisions.

From the viewpoint of traffic safety, failure to identify a drug can have serious consequences if it equates with failure to recognize impairment, and the misses require closer examination of the specific drugs that were missed. The 14 cases where all drugs were missed are listed below. Since five of these arrestees had used multiple substances, a total of 20 drugs were detected.

	Al		_	s Missed rrestees
Narcotic analgesics				_
Morphine				
Codeine	• •	••	•••	1
Stimulants				
Cocaine				5
Methamphetamine		•••	• •	1
Marijuana		• •		5
Depressants				
Barbiturate				1
Benzodiazepine				1
Carisoprodol/Meprobamate				_
Chlorpheniramine				
Meprobamate	•••		•••	1
Other				
		· • •		1

Again, cocaine and marijuana appear most frequently. It is not possible to establish the reasons for misses retrospectively, but misses of cocaine and marijuana are not unexpected. Unless a large amount of stimulant has been ingested, the signs and symptoms typically are less obvious than the symptoms of other categories and can be very difficult to recognize. Cocaine is a fast-acting substance, and observable signs of use may be apparent at roadside but diminish significantly by the time of evaluation. The half-life of cocaine is approximately 90 minutes, but its metabolite, benzoylecgonine (BE), can be detected in urine for 24 - 48 (possibly 72) hours, depending on amount ingested. Thus, it is possible for the laboratory to detect BE from cocaine, which was ingested at some time in the recent past, even though the suspect was not impaired at the time of the evaluation.

Similarly, the marijuana metabolite appears and can be detected in urine for daysto-weeks, depending on amount and chronicity of use. Because a specimen may test positive at a time when the suspect is not under the influence of marijuana, a DRE evaluation is crucial. Importantly, unless a marijuana positive from the laboratory is corroborated with evidence of impairment at the time of the evaluation, it does <u>not</u> speak to the question of drug influence.

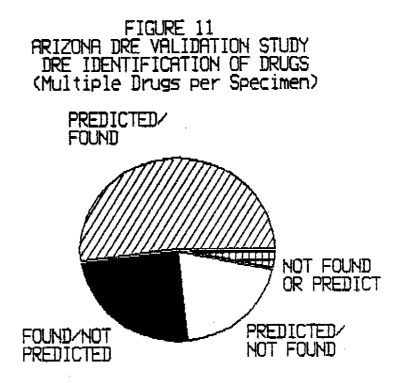
In summary, misses can occur if a DRE fails to correctly observe, record, and interpret the signs and symptoms displayed by a suspect. They will occur if the parent drug has been eliminated from the body, but a metabolite, which is not itself psychoactive, remains in the urine. They will occur if one substance produces severe symptoms, as PCP does, which entirely mask the symptoms of other drugs. Also, although two or more drugs may have been used, differences in amounts used and each drug's time course may be such that not all substances yield signs and symptoms at the time of the evaluation.

Although a true miss and the release of an impaired driver carries the greatest potential for harm, citizens are likely to be understandably distressed by false positive errors. In the PPD data, the DREs believed a drug was present 42 times when no drug was found in the specimen (Table 5, Figure 11). The drug categories, which the DRE believed to be influencing the suspects, are summarized below:

	False Positive (number)
Single Category	
Stimulant	12
Marijuana	· 7
Depressant	5
Phencyclidine	1
Inhalant	1
Narcotic Analgesic	_1
-	27
<u>Two or More Categories</u>	
Marijuana/Stimulant	6
Stimulant/Depressant	4
Stimulant/Narcotic Analgesic	3
Marijuana/Phencyclidine	1
Depressant/Inhalant	1
•	15

TABLE 7ARIZONA DRE VALIDATION STUDYDRE Correct Identifications and Misses, by DrugFor 668 Drug Detections in 416 Specimens

		Num	ber		
	DETECTIONS	CORR		MISSES	
		<u>IDENTIFI</u>	CATIONS		
		<u>Number</u>	Percent		
Marijuana	165	149	90	16	
<u>Stimulants</u>					
Cocaine	115	104	90	11	
Amphetamine	18	17	94	1	
Methamphetamine	69	66	96	3	
Depressants					
Barbiturate	35	33	94	2	
Benzodiazepines	108	106	98	2	
Narcotic Analgesics				`	
Morphine	71	67	94	4	
Codeine	65	62	95	3	
Phencyclidine	_22	_21	<u>95</u> 94	1	
Totals	<u>_22</u> 668	625	94	_ <u>1</u> 43	
Other drugs	145				
Total:					
Drugs detected	813				



- -1

Ten of the arrestees admitted using a prescription drug, and one was in possession of marijuana. None admitted using an illicit substance, and most denied any drug use whatsoever. Stimulants and marijuana appeared most frequently as false positives, as they did for misses.

A more exhaustive analysis of misses and false positives, which is beyond the scope of this project, is recommended. The records now residing in the data base, together with the DIE narratives, will support an analysis of each component of the evaluation. The specific objective would be to examine <u>by drug</u> the specific signs and symptoms, suspects' admissions or denials, and drug possession for each miss and false positive. The relationship of misses and false positives to the time course of each drug, as well as to gender and age characteristics of the suspects, may prove to be variables which predict the errors. If specific signs, symptoms, combinations, and conditions are found to be reliably related to misses and false positives, that information can be incorporated into training and guidelines.

F. Signs and Symptoms and Drug Identification

7

The standardized evaluation enables a trained officer:

- 1) to determine whether a suspect is impaired;
- 2) to determine whether observed impairment is drug-related; and
- 3) to identify the category or categories of drug(s).

As a basis for that three-level opinion, DREs perform the 12-step evaluation in a prescribed, systematic manner and then integrate all of the obtained information. Diverse observations and measures are made during the evaluation, and the relative contribution of the various signs and symptoms to DREs' opinions has not been determined. The following questions are illustrative but not exhaustive of appropriate inquiry:

Does each component of the evaluation (FSTs, eye examination, vital signs, etc.) contribute equally to the DRE's opinion? If not, which is more/less useful?

Does the value of a particular component (or observation) differ by drug or drug combination?

Does the validity and reliability of the method require all components of the evaluation under all circumstances and for all suspected drugs?

When a larger data set becomes available, these questions will be broadly addressed with appropriate and exhaustive statistical analysis. For the present, a data set of 500 cases supports the examination of certain key variables.

1. <u>Eye Signs</u>

The DREs rely on information obtained by examination of the eyes. Among other signs, they look at pupil diameter under various light conditions. For this study, the pupil diameter variable has been analyzed with two different data sets. First, a restricted set of cases, meeting the following criteria, was summarized:

- A single drug was detected in the specimen;
- The detected drug was cocaine, methamphetamine, or morphine; and
- The DRE identified the drug.

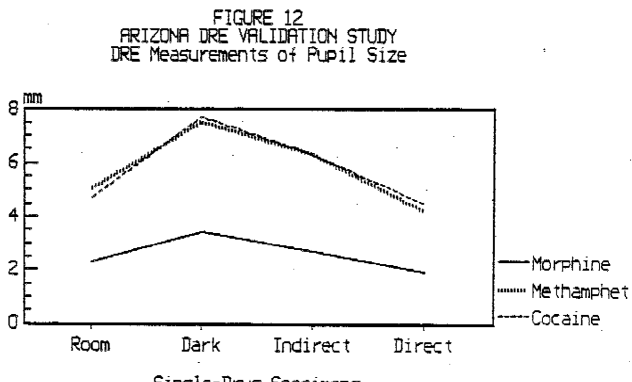
The analysis was limited to cases in which a single drug was detected in the specimen in order to obtain a clear picture of pupillary response to a drug without the possible influence of any other substance, and was further limited to those cases in which the DRE identified the drug. The narcotic analgesic-stimulant comparison was selected because the two drug categories are known to exert opposing effects on pupil size. With these restrictions, the analysis directly addresses the question of whether the magnitude of differences in pupil diameter, as observed by a DRE, was great enough to contribute to drug identification.

A <u>t</u> statistic was calculated for the difference in the darkness condition between observed pupil sizes of suspects under the influence of morphine or cocaine. The mean pupil sizes graphed in Figure 12, together with a <u>t</u> of -6.58 (21 df, p < .01), indicate that the DREs' observations of suspects' pupil sizes were important contributors to drug identification.

A second question focuses on the robustness of pupil measurement in the presence of several drugs since, as can be seen in Table 6, multiple drugs were more common than a single drug. This question has been examined with data for cocaine and morphine. Figure 13 graphs <u>all</u> cases in which either drug was detected, excluding the 29 specimens containing both drugs and also excluding cases with misses and false positives. The data restrictions permit a comparison of observed pupil sizes of suspects who were under the influence of either cocaine or morphine (but with other drugs present) when the DRE identified all drugs present. Again, the diameter of suspects' pupils in the darkness condition discriminated between the two drugs (<u>t</u> -3.97, 114 df, p < .01).

These data confirm that changes in pupil diameter in darkness reliably identify the two drug categories, narcotic analgesics and stimulants. A more extensive analysis is needed to examine the contribution of changes in pupil size and responsivity under other conditions and for other drug categories.

Table 8 summarizes other eye signs for all specimens in which each drug was found. Since the table includes multi-drug as well as single drug specimens, the



Single-Drug Specimens

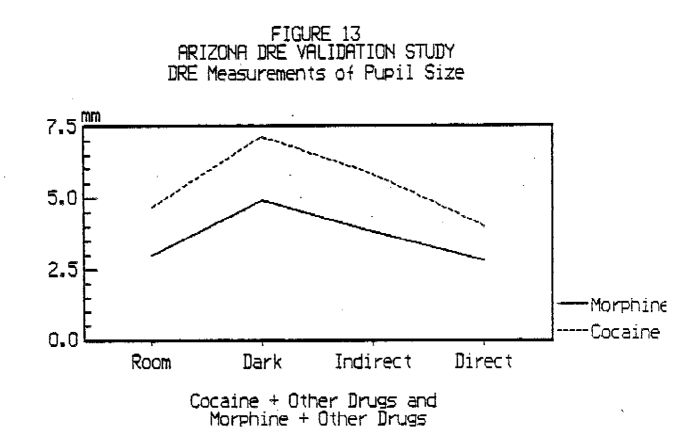


TABLE 8

ARIZONA DRE VALIDATION STUDY Eye Signs Observed during Drug Influence Evaluations Observations (Number, Percent) by Drug Group

EYE SIGNS (See key)		1	2	3	4	5	6	7	8	9	10	11	12
PCP	no. %	<u>16</u> <u>73</u>	18 82	0	<u>12</u> 55	4 18	11 50	<u>5</u> 23	5 23	<u>18</u> 82	<u>18</u> 82	14 64	<u>15</u> <u>68</u>
Morphine	no.	28	56	2	12	14	<u>49</u>	8	<u>39</u>	27	26	29	29
	%	39	79	3	17	20	69	11	55	38	37	41	41
Cocaine	по.	51	79	3	12	<u>29</u>	58	<u>31</u>	44	52	51	52	53
	%	44	69	2.6	10	<u>25</u>	50	27	38	45	44	45	46
Mari.	по.	104	127	1	28	33	61	<u>67</u>	54	97	95	<u>109</u>	108
	%	63	77	0.6	17	20	37	71	33	59	58	<u>66</u>	65
Barbit.	no. %	27 77	29 83	0	<u>15</u> 43	<u>11</u> <u>31</u>	<u>22</u> <u>63</u>	4	<u>14</u> 40	<u>27</u> 77	<u>27</u> 77	<u>28</u> 80	27 77
Benzodiaz.	по.	<u>75</u>	92	4	<u>30</u>	23	<u>74</u>	22	<u>45</u>	<u>70</u>	<u>69</u>	<u>76</u>	<u>76</u>
	%	<u>69</u>	85 -	3.7	28	21	<u>69</u>	20	<u>42</u>	65	64	<u>70</u>	<u>70</u>
Methamphet.	no.	40	43	1	5	<u>24</u>	31	18	31	33	33	39	39
& Amphet.	%	46	49		6	28	36	21	36	38	38	45	45

% = percent of arrestees with the sign whose specimen was positive for the drug Underlined/Bold = drugs with ranks 1, 2, or 3 for each sign

<u>Column</u> 1	<u>Eve Sign</u> HGN
2	Lack of convergence
3	Does not follow stimulus
4	Vertical nystagmus
5	Hippus
6	Droopy eyelids
7	Rebound dilation
8	Slow reaction to light
9	Lack of smooth pursuit, left
10	Lack of smooth pursuit, right
11	HGN at maximum, left
12	HGN at maximum, right
1	

data cannot be used to examine the validity of separate eye signs. An analysis of signs and symptoms when two or more active drugs are present is a complex problem and is beyond the scope of this project. The Table 8 data are presented solely to demonstrate the patterns and trends associated with the various drug categories. As can be seen in the table, "lack of convergence" was recorded for more than half the suspects for all drugs. Thus, it contributes little to the discrimination of any specific drug. Similarly, the value of "not able to follow the stimulus" seems to be limited since it was recorded only 11 times. The other signs show clear-cut patterns despite the presence of multiple drugs in many of the specimens.

The underlined cells in Table 8 indicate ranks 1, 2, and 3 for each sign. To illustrate, "HGN present" is identified in the table as Eye Sign 1 (first column). Note that it was observed in 77% of the barbiturate cases, 73% of the PCP cases, and 69% of the benzodiazepine cases. The preponderance of underlined cells indicate that eye signs are strong predictors for PCP and depressants. Droopy eyelids are associated with morphine, and rebound dilation is associated with marijuana. Fewer underlined cells indicate that these eye signs are less useful for stimulants.

2. Vital Signs

DREs measure a suspect's blood pressure (one time) and pulse rate (three times) during an evaluation. The range of normal values for vital signs is moderately wide and these indices vary as a function of disease and other between-person physiological differences. For these reasons, blood pressure and pulse rate as independent signs and are not expected to have the diagnostic specificity for drugs of the all-or-none phenomena such as horizontal gaze nystagmus (HGN). They are, nonetheless, important cues if they reliably corroborate other observations. A striking disparity, such as depressed vital signs and other observations consistent with PCP, would be cause for further examination.

Table 9 summarizes the blood pressure and pulse rate data for the cases in which the DRE identified a single drug and the laboratory analysis of the specimen confirmed the opinion. Given the small number of cases which meet these strict criteria together with the variability of the measures, the between-drug differences do not reach statistical significance. Although the data in Table 9 are of interest, they should be interpreted cautiously pending replication.

The mean systolic blood pressure for PCP users was 141 mmHg (Table 9). For other drugs, note that the mean values do not exceed the upper limit of the 140/90 normal blood pressure range. The mean blood pressure for suspects under the influence of methamphetamine and PCP was relatively high, as expected. The mean blood pressure with morphine also was elevated in comparison to other

-,					. <u> </u>						
		BLOOD PRESSURE (mmHg)			<u>PULSE RATES</u> (bpm)						
		<u>\$Y</u>	S	DIAS			1	2			3
	<u>n</u>	x	<u></u>	<u>x</u>	<u>7</u>	<u>x</u>	<u> </u>	x	σ	x	<u></u>
Barbiturate	7	124	11	85	9	83	20	84	17	88	18
Benzodiazepine	12	123	15	83 1	7	100	21	101	19	97	20
Cocaine	18	126	20	77 1	5	97	17	97	18	98	16
Marijuana	44	132	18	82 1	5	92	17	94	18	90	16
Methamphetamine	24	133	1 9	85 1	4	100	19	101	20	99	19
Morphine	8	135	20	81 1	3	93	20	99	17	99	20
PCP	5	141	24	87	4	116	27	101	25	116	6

TABLE 9 ARIZONA DRE VALIDATION STUDY Mean Blood Pressure and Pulse Rates * As Measured During Drug Influence Evaluations

* 1 Single drug was detected in specimens <u>and</u> was identified by the DRE without misses or false positives.

2

categories; this unexpected finding may be more instructive about the age and health status of heroin users than about drug effects per se. The finding must be considered highly tentative for the present.

Higher pulse rates (bpm) were recorded with methamphetamine and PCP and also with benzodiazepines. The latter also is an unexpected observation. It is possible, but entirely speculative, to note that it may also reflect arrestee characteristics.

3. Time Estimates

As suspects stand with eyes closed, arms at their side, and head tipped back, they are instructed to estimate a 30 second time interval. Restricting the analysis to cases with a single drug predicted and found, the mean estimates for each drug category appear below.

	<u>Estimate</u> <u>mean</u>	<u>s of 30 sec.</u> <u>std. dev.</u>	
Barbiturates	38	21	50% greater than 30 sec.
Benzodiazepines	38	20	64% greater than 30 sec.
Marijuana	26	12	69% less than 30 sec.
Morphine	27	8	67% less than 30 sec.
Cocaine	22	7	80% less than 30 sec.
PCP	20	7	All less than 30 sec.
Methamphetamine	18	7	92% less than 30 sec.

As expected, depressants tend to lengthen the time estimate and stimulants to shorten it. The estimate appears to be a strong predictor for cocaine, PCP, and methamphetamine. Although the variability in some categories weakens the sign in the individual case, in the context of other symptoms, the time estimates can be expected to serve the DRE well.

G. Arrestees' Drug Choices

Suspects sometimes acknowledge that they have used a drug or drugs. The following table summarizes: (1) arrestees' admissions; (2) in comparison to the number of times the substances were found in suspects' possession; and (3) the positive toxicologies.

	(1) Arrestee <u>Admissions</u>	(2) Drugs Found <u>On Suspect</u>	(3) Positive <u>Specimens</u>
Narcotic	126	19	136 Morphine, Codeine
Depressant .	122	22	143 Barbiturates, Diazepam
Marijuana	97	46	165 Marijuana
Stimulants	78	21	202 Amphetamine, Methamphetamine, Cocaine
PCP	8	1	22 PCP
Inhalant		2	4 Toluene

The high rate of narcotics admissions can be attributed to the addicts' prior experiences in the criminal justice system and their realization that track marks and constricted pupils are uniquely identifying signs. In contrast, marijuana and stimulant users, who may not have been arrested previously, are less likely to understand that the standardized examination enables the DRE to detect their drug use.

Typically, an admission occurs at the conclusion of the evaluation when the DRE has formed an opinion and confronts the suspect about his drug use. The suspect's statements are considered as part of the total evidence, but the DRE is aware that they may be true, partially true, or entirely misleading, and his opinion does not necessarily match the suspect's admission. In these data, when the suspect admitted use of a drug, the DRE identified the drug <u>and</u> it was found in the specimen for approximately 90% of the admissions (range by drug category = 85% to 100%).

VII. DISCUSSION AND CONCLUSIONS

The DRE methodology mandates both a standardized evaluation and the analysis of a specimen. Together, the evaluation and the toxicological analysis create a

balance, which is designed to identify impaired suspects (minimize misses), and equally important, to recognize unimpaired suspects (minimize false positives).

The findings from this study of a set of 500 DIE and SER records provide support for the validity of the methodology. There were few positive DRE opinions which were unsupported by laboratory analysis. The number of false positive opinions and the number of complete misses were low. An accuracy rate of approximately 85% is in agreement with earlier studies.

Analysis of the study records indicates that certain signs and symptoms (pupil size, field sobriety tests, time estimates) are strong indicators of specific drugs. Other signs and symptoms appear to be less strongly linked to a particular drug. Redundant and non-specific symptoms neither enhance nor detract from DRE accuracy, but if careful analysis of evaluation records leads to their identification, it is possible that the evaluation procedure can be simplified.

The DIE and SER records provide insight into the DUID population of Phoenix and their drugs of choice, and into the validity of the DRE methodology. As subsets of the data were examined, however, the numbers became so small as to lack the statistical power to answer questions about specific variables or the interaction of variables. For that reason, the reported relationship between toxicology findings and signs and symptoms are somewhat preliminary in nature. They serve to demonstrate the analytical power of the data base software and the kind of information that can be gleaned from drug evaluation and toxicology records. A number of longer range objectives will be realized as more data become available. In particular, the development of a composite symptom profile for each drug category, validated by analysis of DIE forms and toxicology records, will be undertaken when the number of records support the necessary analyses.

The substances found in this sample of arrestees were largely illegal drugs, although prescription drugs which have a high abuse potential were also found. Although there is a large number of drugs with a potential for affecting the central nervous system, only a limited number of different drugs were actually found in these arrestees. Note that antihistamines and tricyclic antidepressants were rarely a possible factor in causing impairment.

The AZ-DPS Laboratory's analytical protocol detected and confirmed most drugs of interest in driving impairment cases in Arizona. Occasionally, it was necessary to screen for miscellaneous substances (e.g., carisoprodol) by a supplemental secondary screening procedure other than the immunoassay battery. Omitting the secondary screening would have resulted in a lower corroboration rate for DRE opinions concerning narcotic analgesics and depressants, but the merits of the secondary screening must be weighed against the cost to laboratory resources. A comparison of data obtained during this study with data reported by the U.S. Department of Justice (16) is relevant to assessing study findings. During the third quarter of 1992, urine samples were obtained from booked arrestees in 24 drug-use forecasting (DUF) sites. The following rates of "positive for any drug" were reported for Phoenix:

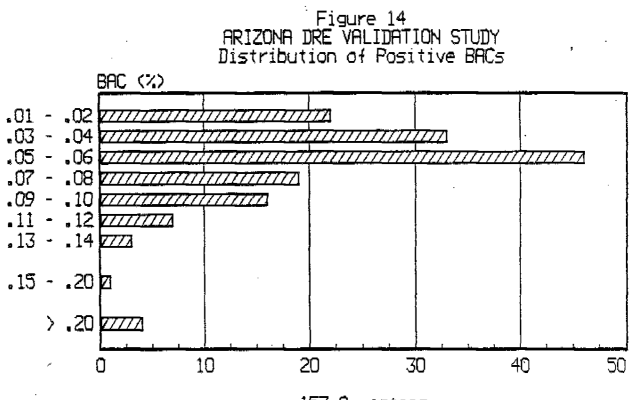
•	<u>% Positive</u>
Juvenile Male Arrestees/Detainees	29
Male Booked Arrestees	54
Female Booked Arrestees	66

The number of men in Phoenix who were drug positive ranks 19th among 24 sites; i.e., <u>fewer</u> men were found drug positive in only five other cities. The rank for women is higher (13th).

In both the DUF and DRE data, marijuana and cocaine are top-ranked drugs-ofchoice, confirming that these two substances are popular with both the general population of drug users and with drug users who drive. The comparisons suggest that, as expected, drug use by traffic offenders reflects drug use in the general population and that traffic officers arrest users of the most common drugs in a community.

Importantly, most of the drivers in this study could not have been arrested and prosecuted without the evidence of impairment obtained from the DRE evaluation and the corroboration by analysis of urine or blood. Figure 14 plots the distribution of positive BrACs in the sample of drug-impaired drivers. Slightly less than one-third of the arrestees had consumed alcohol, and only 5% of the positive BrACs were 0.10% or higher. The suspects with BrACs at and above 0.10%, including four above 0.20%, would have been charged with DUI with or without recognition of their drug impairment. Without the drug influence evaluation, however, the majority of these impaired drivers would not have been held or charged with an offense.

The PPD DREs have been responsible for the temporary removal of at least 378 drug-impaired drivers from Phoenix roadways. At a minimum, those drivers were prevented on at least one occasion from driving in a condition with the potential for harm to themselves and others. Whether the program exerts a longer term deterrent effect upon the arrested drivers, whether it influences the general driving population to avoid driving while impaired, and what the impact of such deterrent effects might be on traffic safety in general are questions which remain to be answered.



153 Arrestees

The major conclusions of this study are:

- The DRE program is a valid method for identifying and classifying drugimpaired drivers.
- Certified DREs recognize drug-impairment and identify the drug(s), by category, which cause the impairment.
- Observable signs and symptoms are associated with specific drugs.
- Monitoring DRE opinions and laboratory results will facilitate program management.
- The DRE program requires scientifically sound support by the laboratory.

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APPENDIX I

:

ROSTER OF DRES

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STUDY PARTICIPANTS

Phoenix Police Department

Chief Dennis Garrett Sgt. Richard Yost, DRE Coordinator Lt. Joe Klima, Past DRE Coordinator

Off. Larry Babcock Off. Richard Bartlett Off. Mark Beadles Off. C.E. Buddle Off. Mike Campbell Off. Ramsey Campbell Off. A.R. Contreras Off. S. Durham Off. Toby Ehrler Off. Michael Greenfield Off. Timothy Hallahan Off, Vern Hancock Off. Richard Hyde Off. Herbert Jacobs Off. B. L. Kelly Off. Joe Knott Off. William Lee II Off. Doug Marks Off. Jerry McFarland * Off. Frank Milstead Off, Ronald Nagy Sat. Bill Niles Off. Tim Overstake Off. Steve Park Off. Bill Sampson Off. Terry Sills Sgt. Robert Sparks Off. Joel Tranter Off. George Tryon Off. Ed Tuttle Off. James A. Unsworth Off. Robert T. Ward

Study Participants, Other Agencies:

Sgt. Claudia Baca, Arizona DPS Sgt. Robert Hohn, Arizona DPS Off. Gary Horner, Glendale PD Off. S. Twitchell, Scottsdale PD

* Officer McFarland, who became a DRE early in the Arizona program, passed away prior to the time period of the study.

RECENTLY CERTIFIED DRES

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Phoenix Police Department

Off. Michael Adams

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Off. Douglas C. Callicotte

Off. Jeffrey A. Chapman

Off. Mark R. Hafkey

Off, Michael Henderson

Off. Gregory A. liames

Off. Gary L. McCarthy

Off. Lance D. Miller

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Off. Timothy D. Norton

Off. David Pallis

Off. Michael E. Sales

Off. Edward L. Smith

Off. James R. Smith

Off, Harold A. Sprouse

Off. Ross V. Taylor III

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OTHER ARIZONA DRES AND AGENCIES

Arizona Department of Public Safety

Off. Vern Alley, Coordinator (Statewide) Sot, Claudia Baca, Past Coordinator (Statewide) Lt. Robert Halliday, Past Coordinator (Statewide) Sqt. Robert Hohn, Past Coordinator (Statewide) Off. Jerry Oldsen, Past Coordinator (Statewide) Off. Guy Anderson Off, Edward Andersson Off. William Arthur Off. Michael Bonin Off. John Bottoms Off. Bruce Campbell Off. Marty Camacho Off. Gary Ciminski Off. Pete Drummond Off. Mike Crowe Off. Thomas Eaves Off. Brian Eekhoff Off. Jaime Escobedo Off. Wolfgang Evans Sgt. Michael Fane Off. Brett Farrar Off. Regina Georgitso Off, John Gigous Off. Tim Goodwin Off, Jack Hegarty Off. Kevin Jex Off. Jeff King Off. Michael Livingston Off, Daniel Lugo Off. Dale Mace Off. Mike Macias Off. Paul B. Maine Off. Bobby Marquez Off. Jeff Nash Off. Daniel Ortiz Off. Robert Osborn Off. Stephen R. Reutter Off, Randy Roby Off. Dan Slade

Arizona Department of Public Safety - Continued

Off. Ann Stuckey Off. Steve Tritz Off. Robert Ticer Off. Rene Valencia Off. Rick Valencia Off. A. S. Vildusea Off. Johnny Villaneda Off. Johnny Villaneda Off. Bruce Weddle Res. Dennis Duffy Res. Bert A. Stanfield-Pinel Res. Richard Studdard (retired, past coordinator, LAPD)

Apache Junction Police Department

Off. Troy Mullender

Avondale Police Department

Off. Patricia Stinson. Off. M. Reynolds

Buckeye Police Department

Off. Charles V. Griffis

Casa Grande Police Department

Off. Michael Colvin

Chandler Police Department

Off. Kurt Hauser Off. John Porvaznik Off. Mike Slupinski

Gilbert Police Department

Off. Scott Hanson Off. Mike laquinto

Glendale Police Department

Off. Mike Stockton, Coordinator Off. Brent Coombs Off. Gary Horner Off. Brian Lahti Off. Jim Reynolds Off. Mark Smith Off. Brian Wilkins Cpl. Steve R. Willis

Lake Havasu Police Department

Off. Rick Eyestone Off. Eugene Radecki

Maricopa County Adult Probation

Nancy S. O'Brien

Maricopa County Sheriff's Office

Dep. John W. Allen Dep. Leslie Paul White

Mesa Police Department

7

Sgt. Steve Toland, Coordinator Off. Trish Bradley Off. Dan Brown Sgt. Richard Clore Off. Jerry Gissel Off. William Green Off. Royed B. Hollick

Mesa Police Department - Continued

Off. Jay Hutson Off. Brian Kozak Off. Ron Martinez Off. Donald Moss Off. Manny Quinonez Off. Dave Rhodes

Mohave County Sheriff's Office

Dep. Don Bischoff Dep. Robert N. Kuerner Dep. Scott Kuerner

Northern Arizona University Police Department

Off. Bryan D. McKinnon

Paradise Valley Police Department

Off. Vincent Leone

Peoria Police Department

Off. Rich Scrivens Jr. Off. R. J. Smith

Pima County Adult Probation

Linda Gloy

Pima County Sheriff's Office

Dep. Manuel A. Amado Dep. Bill Brantley Dep. William D. Murphy Dep. Christopher Radtke

Scottsdale Police Department

Off. Shawn Twitchell, Coordinator Off. Jeffrey Belford Off. James Butera Off. William Monahan Off. J. Jeffrey Smythe

Sierra Vista Police Department

Off. Robert Randall

Surprise Police Department

Off. Claude Carroll

Tempe Police Department

Sgt. Toby Dyas, Coordinator Off. Gerald Adams Off. Randall Fougner Off. Bob Gage Off. Robert Johnson Off. Dave Lind Off. Richard Tabor Off. Ed Wells

Tucson Police Department

Sgt. John Patla, Coordinator Off. Nicolaas Aussems Off. Ramon Batista Off. George Eppley Off. Richmond E. Holley III Off. Robert Jenkins Off. Clayton Kidd Off. Clayton Kidd Off. Wayne Martinez Off. Timothy Milbourn Off. James Monaco Off. Mark Napier

Tucson Police Department

Off. Phillip Penta Off. Kathy Pipes Off. Michael Pryor Off. Dennis Qubik Off. Gary Scaramuzzo Off. Carlos Valdez Off. Kathryn Wendling

Wickenburg Police Department

Off. Joe Favazzo

APPENDIX II

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DRE COURT CASES AND HEARINGS

<u>State v. Johnson et al.</u> Cit 90056865, (1992), <u>Frye</u> hearing, Tucson, Arizona. Held: DRE meets Frye test. Special action jurisdiction to Supreme Court, denied. <u>Johnson et al. v. Hon. Rita Jett</u> (Real Party in Interest, City of Tucson) CV-91-0488-SA (1992).

People v. Quinn, 580 N.Y. Supp. 2d 81 (Dist. Ct. 1991); <u>Frye</u> hearing, Dist. Court Suffolk County, New York. Held: DRE meets Frye test (appeal pending).

People v. Hernandez, No. 92M181 (1992); Frye hearing, County Court, Boulder, Colorado. Held: Frye inapplicable; DRE testimony admissible.

<u>State v. Klawitter</u>, CA-93-2092; (1993); <u>Frye</u> hearing, Minneapolis, Minnesota. Held: <u>Frye</u> inapplicable; DRE testimony admissible (pending special action to the Minnesota Supreme Court).

APPENDIX III

2

Drug Influence Evaluation Form Scientific Examination Report Checkprint Template Blood Drug Analysis Form Urine Drug Analysis Form

A

ARIZONA DEPARTMENT OF PUBLIC SAFETY SCIENTIFIC EXAMINATION REPORT

OR NO.

AGENCY Phoenix F.D.

FILE NO.

OFFICER

DATE

NAMES]

ITEMS :

#1. Orine specimen

EXAMINATION REQUESTED :

Drug Screen: narcotic analgesic, CNS depressant

RESULTS :

Item #1 - Analysis of the wrine showed it to contain methadone, propoxyphene, norpropoxyphene, desalkylflurazepam and hydroxyethylflurazepam (metabolites of flurazepam), alpha-hydroxyalprazolam (a metabolite of alprazolam), chlorpheniramine, and diphenhydramine.

CUSTODY OF EVIDENCE

RECEIVED D.P.S. Property

DISPOSITION D.P.S. Property

EXAMINER

Distribution: Department Records, Exeminer/Lab. file. Officer, Prosequitor

DEPARTMENT RECORDS

OPE 802-01950 Per.

	SCIENTIFIC EXAM	<u>T OF PUBLIC SAFETY</u> INATION REPORT	
AGENCY 3	Phoenix P.D.	GR NO.	
FILE NO.			
orrezk ·			
GATE			•
NAME(S)			
ITEMS	â		
#1. (Orine specimen		
EXAMI	VATION REQUESTED :	•	
Drug	Screen - Narcotic Analgesic		
	· ·		
RESUL'	80 .		
	Analysis of the urine show 6-monoacetylmorphine (a me	wed it to contain morphing, codef stabolite of heroin), methadone, plite, cocaine and methylecgoning	
custody of e received	Toxicology Coldroom	EXAMINER	
DESPOSITION	D.P.S. Property		

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DEPARTMENT RECORDS

				PAG	36	OF
					NUMBER:	
				ΕV	ALUATOR:	······································
			1001	CO	NTROL #:	
DRUG INFLUENC		LUAI	IUN	BO	DKING #:	
ARRESTEE'S NAME (Last, First, Mt)		AGE SEX	RACE A	RRESTING	S OFFICER (Name	e, Badge, District)
DATE EXAMINED/TIME/LOCATION	1	EATH RESULT	TS: 🖸 Refused Instrumen			MICAL TEST Doth Tests Urine DBlood Refused
MIRANDA WARNING GIVEN: Yes Given by: D No	What have	you eaten tod	lay? When	?	What have you b How much?	een drinking? Time of last drink?
Time now? When did you last sleep? How long?	Are you si	ck or injured?		C Yes C No	Are you diabatic	<u> </u>
Do you take (asulin ?	Do you hi	we any physic:	al defects?	U Yes	Are you under st doctor/dentist?	se care of a O Yes O No
Are you taking any medication or drugs? Ves No	ATTITUO)É			COORDINATIO	N
SPEECH	BREATH				FACE	
CORRECTIVE LENS: DNone	Eyes:	al 🗋 Bloods	hot OWatery	8lindnes:		L. Eye GEqual DUnequal
PUPIL SIZE: C Equal Unequal (explain)		HGN Preser	ici CINe	Able to fo	How stimulus:	Eyelids:
PULSE & TIME HGN	Right Eye	Left Ey		il Nystagmi	15? OYes ON	ONE LEG STAND:
1 Cate of Smooth Forser			Conver	tignt Eye	Lett Eye	
3 Angle of Onsec				$\underline{}$	\bigcirc	A A I A
BALANCE EYES CLOSED WALK AND TURN	TEST		Cannot keep Starts too soo			
		-h	Stops Walkin		st Nine 2nd Nine	
			Misses Heel-T			L R D D Sways while balancing ,
	حصحص		Steps off Lin Raises Arms			C Uses arms to balance.
			Actual Steps	Taken		Puts foot down.
INTERNAL CLOCK: Describe Turn Estimated as 30 sec.			Cannot do Tesi	r (explain)	Type of Footwear
C Rignt 🛆 Lan	PUPIL SIZE	Room Light	Darkness	Indirect	Direct	NASAL AREA
Draw jines to spots touched	Left Eye Right Eye					ORAL CAVITY
	HIPPUS		REBOUND DI	LATION	Reaction to	Light
		C No	CIYes .	QNo		
		e			LEFT ARM	
ا من الأ		<u>}</u>	<u>)</u>	_		
				L	AV.	<u> </u>
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BLOOD PRESSURE: TEMP		\leq				\sim
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MUSCLE TONE: O Near Normal D Flacid ORigid Comments:		ATT	ACH PHOTO	 5 OF FR	ESH PUNCTU	REMARKS
	low much?	Time	of use?	When	e were the drugs us	ed? (Location)
DATE/TIME OF ARREST	T	IME DRE NO	TIFIED	EVA	L START TIME	TIME COMPLETED
OFFICER'S SIGNATURE			DISTRICT		ID NUMBER	REVIEWED BY
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	EYE REACTION TEST RESULTS
: :Has HGN : :Does NOT Fo : :Tracking NG : :Vertical Nn : :Lack of Con	: Droopy Eyelids : Pupils NOT equal IT equal : Rebound Dilation rstasmus : Hippus
PUPIL SIZES	ᆂᆧᅸᇏᆕᅒᇭᆍᆋᆑᅸᆂᆂᇌᆕᆕᇻᆂᆂᇑᇊᅏᄟᇳᅖᅓᅸᇏᇑᇊᆕᆂᅶᅶᇍᇍᆋᆋᅸᆂᇍᆃᆂᄣ ᆃ
roric 31253 LEFT oom Light . arkness . ndirect Light . irect Light .	# EYE MOVEMENTS RIGHT # * Lacks Smooth Pursuit : : : * Max. Deviation HGN : : : : * Angle of Onset
╧╬╤┓ᆍ┕╍╩╬╬╦╕╕┲┲┟╬╤╕┲╶	DRUG USE REPORT
DRUGS FOUND : :COCAINE : :STIMULANTS. : PHENCYCLIDINE : :HALLUCINOGENS : :CANNABIS : :CANNABIS : :ALCOHOL : :DEPRESSANTS : NARCOTICS : NOT IDENTIFIED : :DTHER Test Type: : :Uri : :Bid : :Ref	
Positive : P.C.P : Opiates-Moret : Codeine : Cocaine : Cocaine : Marijuana : Barbiturates. : Valium : Methaqualone : Amehetamine : Other Other:	hine : : : :

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NOTES

BLOOD DRUG ANALYSIS

Received:	:	DR #
Returned:		ltem #
DRE		I NON-DRE

Notes: Evidence Description

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Date

Analyst

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POS	NEG		CONFIRMED	NOT CONFIRMED				
		Cocaine/Metabolite						
		Methamphetamine	· 🔲					
		Opiates						
		Barbiturates						
		Benzodiazepines						
		······································	. 🗖					
		>						
Analyst	/Date		Analyst/Date					
ROUTI	ng: [Further Screening	🔲 cc/мs					

Secondary Screening:

Analyst/Date___

Confirmation Analysis:

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Analyst/Date:_____

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NOTES

URINE DRUG ANALYSIS

Received:	DR #	
Returned:	Item #	
D DRE	DRE CER	NON-DRE
Notes: Evidence Description	Date	
	Analyst	
RIA	G	C/MS
POS NEG	CONFIRMED	NOT CONFIRMED
Cannabinoids		
Cocaine/Metab	olite	
Methamphetam	uine 🔲	
D D Opiates		
Barbiturates	Ū	
Benzodiazepin	.	
Analyst/Date	Analyst/Date	
	ning GC/MS	

Secondary Screening:

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Analyst/Date_

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GC/MS Confirmation Analysis:

THCA-TMS

 $\Box = \Delta^{2} \text{ THCA-TMS}$ $\Box = \Delta^{3} \text{ THCA-TMS} (IS)$

- BENZO-TMS
- desaikyiflurazepam-TMS
- desmethyldiazepam-TMS
- 🖸 🛛 oxazepam di-TMS
- bromazepam-TMS (IS)
- 🗆 👘 lorazepam di-TMS
- temazepam-TMS
- OH-ethylflurazepam-TMS
- -OH-alprazolam-TMS
- 🗇 –-OH-triazolam-TMS

<u>Other</u>

Analyst/Date_

BZE-TMS

- C cocaine
- C methylecgonine-TMS
- benzoylecgonine-TMS
- scopolamine-TMS (IS)

OPSIM-TMS

- dihydrocodeine-TMS
- codeine-TMS
- hydrocodone-TMS
- □ morphine di-TMS .
- C oxycodone-TMS
- □ 6-monoacerylmorphine-TMS
- alorphine di-TMS (IS)

APPENDIX IV

DATA BASE DIRECTORY OF RECORDS

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AGE NO. 03/18/94

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DIRECTORY OF DATABASE 03/18/94

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4 100004		PPD	4	2534	-	01/23/89		M	2	6
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$\begin{array}{c} 11 & 100011 \\ 12 & 100012 \end{array}$	AZ	PPD	12		З	02/24/89		F	2	6
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23 100023	AZ	PPD	24	3930	1			M	5	4
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26 100026	AZ	PPD	24		7	04/16/89		M M	22	6 6
27 100027	AZ	PPB	27		10			M	ŝ	6
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31 100031 32 100032	AZ	PPD	32		1	06/25/89	21	Μ	2	é
33 100033	AZ	PPD	33		2	3 06/26/89		F	2	
34 100034	AZ	PPD	34	_	1			M	5	
35 100035	AZ	PPD		5 2534	(5 06/07/89		M	2	
36 100036 -	AZ	PPD		5 4149		3 05/25/39			2	
37 100037	AZ	PPD		7 2534		7 05/28/89 1 07/14/89			22	
33 100038	AZ	PPD	30	3 4009 9 4228		1 07/14/8: 1 07/08/8:			-	2 6
39 100039	AZ	ppd Ppd		9 4443		1 07/06/85			2	6
40 100040 41 100041	AZ AZ	PPD PPD		1 3759		1 08/31/8			1414 1414	2 6
41 100041 42 100042	AZ	PPD		2 3807	:	2 08/25/8	9 20			5 6
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44 100044	AZ	PPD		4 4443		2 08/13/5		M	- - 	6
45 100045	AZ	PPD		5 2299		1 09/16/8		F	, , ,	26
46 100046	AZ	PPD	4			2 09/27/8		F M		36 26
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RECORD NUMBER			AGENCY	AGENCY COUNT	DRE OFFICER	DRE COUNT	ARREST DATE	AGE	SEX	A C =E	0 R =K
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49	100049	AZ	PPD	49	3304		09/10/89	29	F	2	6
50	100050	AZ	PPD	50	4339	1	09/07/89	51	M	2 2	Ģ
51	100051	AZ	PPD	51	3807	3	09/05/89	35 33	M M	2	6 6
52	100052	AZ	PPD PPD	52	2534 4339	2	06/14/90	25	M	5	é
53	100053	AZ	PPD	53 54	4337	1	06/27/90	63	F	2	р 6
94 55	100054 100055	AZ AZ	260 260	55	3307	4	06/30/90	28	M	2	6
55 56	100056		- FD - FPD	56	3759	4	05/25/90	36	F	2	Ä
30 57	100056	AZ	PPD		4192	2	05/25/90	36	M	2	ā
58	100058	AZ	PPD		3807	24	05/24/90	21	F	2	5
00 39	100059	AZ	FFD	59	4149	4	05/22/90	31	M	2	4
60 60	100040	AZ	PPD	60	4037	11	05/19/90	20	M	2	ė
61	100061	AZ	FFD	61	4228	2	05/15/90	27	M	2	6
62	100062	AZ	PPD	62	3398	3	05/14/90	28	М	2	6
	100063	AZ	PPD	63	4339	З	05/04/90	18	M	2	6
64	100064	AZ	PPD	64	3385	4	05/03/90	25	F	2	6
65	100065	AZ	Dad	65	4339	4	05/02/90	29	M	2	6
66	100066	AZ	PPD	66	3807	5	05/02/90	25	М	2	6
67	100067	AZ	PPD	67	4017	_ 1	08/31/90	4 Q	F	2	6
68	100068	AZ	PPD	63	2985	1	08/29/90	22	F	2	6
59	100069	AZ	PPD	69	2424	. 1	08/22/90	45	F	3	6
70	100070	AZ	PPD	70	2885	2	08/23/90		M	2	6
71	100071	AZ	PPD	71	4293	2	08/23/90	25	M	2	6
72	100072	AZ	PPD	72	4017	2 3	08/24/90	16 31	r M	22	4 6
73	100073 100074	· AZ	PPD PPD	73	4293 3996		08/20/90	29	M	2	6
74 75	100075	AZ AZ	PPD	75	3437	1	08/17/90	27	M	$\frac{4}{2}$	6
7.5	100076	AZ	PPD	76	3304	ź	08/14/90	$\tilde{40}$	M	ŝ	ĕ
77	100077	AŻ	PPD	77	2534	, 9	08/11/90	18	м	5	6
78	100078	AZ	PPD	78	2424	2	08/11/90	34	M	2	- 6
79	100079	AZ	PPD	79	4293	4	08/08/90	37	M	5	6
20	100080	AZ	PPD	80	3901	1	08/07/90	37	F	- 2	6
31	100081	ΑZ	PPD	61	2534	10		- 28	М	2	6
82	100082	AZ	PPD	32	3807	6	03/22/90	22	M	З	\diamond
83	100083	ΑZ	PPD	83	4037	12	03/20/90	34	M	З	4
84	•	AZ	PPD	84	4149	5	03/16/90	17	M	2	6
95		AZ	PPD	95.		4	03/10/90	20	M	2	4
	100086	AZ	FPB	86	2534		03/10/90		M	22	2
	100087	AZ	PPD Bod	87			07/08/90 03/04/90	32 22	E M	- 1	6 8
සිට දෙක්		AZ	pop	38	4339 4443		03/04/90	21		4 2	6 6
25 03		AI AI	299 295	20 29			03/04/30	45		3	⊖ A
	100070	64 - 2 -	250 292				02/01/90			2	5 5
7.		AZ	PPD		4339		03/01/90			$\tilde{2}$	ė
	100092	AZ	PPD		4192		03/06/90			2	ě
	100094	AZ	FPD				03/30/90			3	6
	100095	AZ	PPD	95	3528	1				2	-5
	100096	AI	FPD	96		6	02/14/90	99	Μ	2	6

DIRECTORY OF DATABASE 03/18/94

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4

PAGE NO. 03/18/94

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RECORD NUMBER	CONTROL NUMBER	STATE	AGENCY	AGENCY COUNT	DRE OFFICER	DRE COUNT	ARREST DATE	AGE	SEX	A C ≖E	O R ≖K
											-
			* **	~~	4007	10	00/10/00	90	м	~	,
	100097	AZ	PPD PPD		4037 4037		03/10/90 02/11/90	99 17	M	2 2	- 6 - 6
98 99	100098 100099	AZ AZ	PPD		4037		02/10/90	20	M	â	- 6
	100100	AZ	PPD		4228		02/08/90	ŝŝ	F	$\overline{2}$	4
	100101	AZ	PPD		4149		04/07/90	20	M	2	6
	100102	AZ	PPD				02/06/90		F	$\overline{2}$	6
	100103	AZ	PPD				02/02/90		F	З	6
104	100104	AZ	PPD	104	2705	3	01/15/90	24	М	2	6
105	100105	AŻ	PPD		4228	4	01/14/90		М	2	6
106	100106	AZ	PPD		4009	2	01/12/90	19	M	2	6
	100107	AZ	999		2299		01/12/90	21	м	5	6
	100108	AZ	PPD		3385		01/05/90	34	F	2	6
	100109	AZ	PPD		4443		01/02/90	43	F	2	6
	100110	AZ	PPD		3378		04/30/90		۴	. 2 . 5	6
111	100111	AZ	PPD PPD		3759 4228		04/28/90	27	M M	. J 3	6 6
	100112 100113	AZ AZ	PPD		4149		04/26/90	30	M	5	
	100113	AŽ	PPD		3759		04/20/93	27	M	š	6
• •	100115		PPD	115		_	04/15/90	29	M	6	6
	100116	AZ	PPD		4037		04/08/90	23	M	3	6
117		AZ	PPD		3186		04/14/90		M	2	6
113	100118	AZ	PPD	113	3385	· 8	04/07/90	13	м	2	6
119	100119	AZ	PPD	119	2705	4	04/01/90		М	- 2	6
120	100120	AZ	PPD	120	2534	13	04/06/90	29	М	2	6
121	100121	AZ	PPD		3996	2	08/07/90	· -	F	2	6
122	100122	AZ	PPD		3125	Ţ	08/06/90		M	25	6
123	100123	AZ	PPD		4017	3		17	M	5	6
124	100124	AZ	PPD		4293		08/02/90	23	M	2 2	6
125		AZ	59D			2		67 00	M	2	6
126 . 127	100126	AZ AZ	PPD PPD	120	3437 3901	32	08/01/90	33 35	M F	22	6 6
. 127 128	100127 100128	AZ	660 660		3125		09/23/90	31		2	6
128	100129	AZ	PPD		4017	4	09/28/90	43		2	6
130	100130	AZ	PPD		4293	6	07/20/90	13	M	$\overline{2}$	6
131	100131	AZ	PPD	131		4	09/18/90	22	м	2	6
132		AZ	PPD		3701	1	09709790	42	M	2	6
133	100133	AŻ	PPD	133	3525	1	09/09/90	42	м	- 2	6
134	100134	AZ	PPD		3125		10/30/90	41		6	6
	100135	AZ	PPD		3807		10/30/90			6	6
	100136	AŻ	PPD		3525		10/25/90			2	6
	100137	AZ	PPD		2534		10/24/90			2	é
	100138	AZ	PPD COD		3398	-	10/21/90			2	6
	100139	AZ	PPD PPD		4037 3525	17				2	6
	100140 100141	AZ AZ	PPD PPD		3525 3791	32	10/06/90 10/03/90			22	6
	100142	AZ	PPD		4443		11/30/90			2	6 3
	100143	AZ	PPD		3901	3				$\hat{\bar{z}}$	6
	100144	AZ	PPD		4443		11/17/90			2	6

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DIRECTORY OF DATABASE 03/18/94

RECORD NUMBER	CONTROL NUMBER		AGENCY	AGENCY COUNT	DRE OFFICER	DRE COUNT	ARREST DATE	AGE SEX		0 R ∺K
146	100145 100146	AZ AZ	PPD PPD	146	3996 3701	3	11/16/90 11/13/90	27 M 28 M	10 Q I	4 6
148	100147 100148	AZ AZ	PPD PPD	148	4443 3901	8 4	11/13/90 11/08/90	30 F 28 F	22	1 <u>.</u> 12
150	100149 100150	AZ AZ	PPD .PPD		3385 3701	9 5	11/03/90 12/29/90	27 M 32 F	αN	6 6
151 152	100151 100152	AZ AZ	PPD PPD	151 152	3398 ,3839	7	12/29/90	36 M	22	6 6
153 154	100153 100154	AZ AZ	PPD PPD	153 154	3701 2534		12/20/90	36 M	2 2	6 6
155 156	100135 100156	AZ AZ	PPD PPD	156	4017 3398	5 8	12/17/90	48 M	2 2	6 6
	100137 100158	AZ AZ	ppd Ppd	158	4443 2885	9	12/14/90	19 M	2 2 2	6 6
	100159 100160	AZ AZ	PPD PPD		3807 3437	-	12/04/90	37 M	22	6 6
161 162		AZ AZ	PPD PPD		3701 3528	02	01/28/91	28 M	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	6 6 6
164		· AZ AZ	PPD PPD	· 163	2885 3304 3378	4 9 9	01/31/91 01/29/91 01/27/91	29 M	22	6 6 6
166	100165 100166 100167	AZ AZ AZ	РРD РРD РРD	165	3437 3807	6	01/17/91	48 M	· 2 3	' 6 3
168		AZ	PPD PPD	167	2376 4149	1 9			22	6
170		AZ AZ	PPD PPD	170	4017 4223		01/10/91		2	6 6
172 173	: 100173	AZ AZ	PPD PPD	172	2534 : 4037		03/30/91	22 M	3 2	6 6
174	100175	AZ	PPD PPD		3807	11	03/25/91	. 22 M	23	6 4
177	100176	AZ AZ	990 990 990	176	: 2885 : 3398 : 3525		03/20/91 03/26/91 03/20/91	1 56 M	2 2 2 2 2	1 6 6
179	100178 100179 100180	AZ AZ AZ	PPD PPD	178	3323 3 3839 7 4192	72	03/15/9:	L 22 M	4 3 2	
181		AZ	PPD PPD	180) 3807 . 2885	12		1 26 M	22	6
18: 18:	3 100183 4 100184	AZ AZ	PPD PPD	162 183	2 3 4 37 3 3807	13	7 03/02/9: 3 03/01/9:	1 33 M 1 40 F	2 2	6 6
134	5 100185 5 100186	AZ AZ	PPB PPD	185	1 3398 5 3378	12	02/25/9	1 49 M	2 2	6
18	7 100187 3 100188	AZ AZ	PPD PPD	137	5 2885 7 3437		7 02/14/9 3 02/07/9	1 22 M	25	ι <u>ζ</u> .
13		A2 A7	PPD PPD		3 4017 9 4017		7 02/02/9 3 01/30/9			

PAGE NO. 03/18/94

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5

DIRECTORY OF DATABASE 03/18/94

RECORD NUMBER	CONTROL NUMBER		AGENCY	AGENCY COUNT	DRE OFFICER	DRE COUNT	ARREST DATE	AGE SEX	A 0 . C R =E ≖K
	100193	AZ	PPD		4037	19	11/04/89 11/06/89	34 M 33 M	3 6 2 6
	100174	AZ	PPD		4339	8 20	11/08/89	27 M	36
	100195	AZ	PPD		4037	. 17		43 M	26
	100196	AZ	PPD	195	2534	· 1/ 9	11/30/89	27 F	26
	100197	AZ	PPD		4339		11/26/39	56 M	2 0
	100178	AZ	PPD		3398 4037	21	11/21/89	17 M	5 6
	100199	AZ	PPD	198			11/15/39	16 M	26
	100200	AZ	PPD		4037		11/08/87	26 M	$\frac{2}{3}$ 6
201		AZ	220 220		4149		12/30/89	23 M	56
	100202	AZ			3378	15	12/30/89	31 M	
203	100203	AZ	660 660		4149		12/26/89	40 M	26
	100204	AZ AZ	PPD PPD		2534		12/22/89	19 M	2 6 2 6 2 6
205		AZ	PPD		3930	2	12/21/39	22 M	26
	100204	AZ	PPD		4149	12		27 M	5 6
	100207	AZ	PPD		3398				26
	100208	AZ	229 299		3759	10		46 M	26 24
	100209 100210	AZ	PPD		3528		12/12/89	33 M	26
	100210	AZ	PPD		4037		12/23/89	20 M	26 26
	100212	AZ	PPD		4443		12/27/89	57 M	26 26
	100213	AZ	PPD	212		9	12/09/89	35 M	26
214		AZ	PPD	213		4	12/10/89	24 M	26
	100215	AZ	PPD	214		19	12/07/89		26
	100216	AZ	PPD	215	4037	24	12/06/39	24 M	26
217		AZ	PPD	216	4192	5	12/06/89	19 M	56
	100218	AZ	ppd	217	4192	6	12/05/89		26
	100219	AZ	PPD		2885	8			2 6
	100220	AZ	PPD		3901	6			56
221	100221	AZ	PPD		3398		07/09/90		26
222		AZ	PPD	221		3	-		26
223	100223	AZ	PPD	222			07/16/90		56
224	100224	AZ	PPD		4017	9			26
225		AZ	PPD	224			07/21/90		26
226		AZ	PPD	225		_	07/22/90		26
227		AZ	PPD		3996	4			21 26
228		AZ	PPD	227		6	07/31/90 07/01/90		26
	100229	AZ	PPD	228			07/02/89		26
	100230	AZ	PPD		4192		07/31/89		$\frac{1}{2}$ 6
	100232	AZ	₽₽Ŭ OOD		4443 3304		04/04/89		26
	2 100233	AZ	660 660		3 2534		03/20/89		$\frac{1}{2}$ $\frac{1}{4}$
	3 100234	AZ	PPD PPD		\$ 2334 \$ 4037		03/03/89		2 ě
	100235	AŽ AZ	PPD		5 4149		01/30/89		26
	5 100236		PPD		5 4192		02/24/92		$\frac{1}{2}$
	6 100237	AZ AZ	PPD		7 4192		04/20/93		2 6
	7 100238	AZ AZ	PPD PPD		, 4192 3 4149		03/09/91		$\frac{1}{2}$ $\frac{1}{6}$
	3 100237		PPD		9 3996		5 07/18/9:		26
	9 100240	AZ	PPD PPD		7 3776 D 4228		07/03/91	-	$\frac{1}{2}$ $\frac{3}{4}$
240	0 100241	AZ	FFU	241	· *****	,		_	_ ·

DIRECTORY OF DATABASE 03/13/94

2

PAGE NO. 03/18/94

6

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RECORD NUMBER				AGENCY COUNT	DRE OFFICER	DRE COUNT	ARREST DATE	AGE SEX	A 0 C R ≠E ≠K
				_ .		_	~		a /
	100242	AZ	PPD		3125	-	06/04/91	37 M	36
	100243	AZ	PPD		3759	12	01/10/90	37 M 50 M	2 6 2 4
243	100244	AZ	PPD	243	4228		05/23/90	52 M 28 F	2 6
244	100245	AZ	PPD		4149	15	04/14/90	же г 33 М	$\frac{2}{2}$ 6
245	100246	AZ	PPD		4443 3996	6	04/17/91	24 M	2 6
246	100247 100248	AZ AZ	PPD PPD		4228	ŏ	04/23/91	53 M	26
247 248	100248	AZ	PPD	243	4192	10	04/23/90	22 M	$\frac{1}{2}$ 1
248 249	100250	AZ	PPD	249	4037		05/01/91	49 M	34
250	100251	AZ	PPD	250		7	04/16/91	21 M	22 26
251	100252	AZ	PPD	251		27	04/13/91	24 M	26
	100253	AZ	PPD	252	4293	7	04/09/91	27 M	23
253		32	PPD	253		16	04/09/91	32 M	
	100255	AZ	PPD		3701	6	04/05/91	42 F	26
	100256	AZ	PPD				04/02/91	29 M	26
	100257	AZ	PPD	256		7		56 M	26
257		AZ	PPD		2885	9	05/28/91	23 F 27 M	26
	100259	AZ	PPD		4293	_		31 M 46 M	$\begin{array}{ccc} 2 & 3 \\ 2 & 6 \end{array}$
	100260	AZ	PPD	259		11 10		35 M	25
	100261	AZ	890 000	260	4228 4443		05/11/91	33 M	2 5 2 3
261	100262	AZ	PPD PPD	261		13	05/13/91	33 M	26
262	•	AZ AZ	PPD	263		14		19 F	26
263 264		AZ	PPD		4443	14		19 M	23
265		AZ	PPD		2534	21	05/02/91	21 F	24
266		AZ	PPD		2885	10	05/01/91	21 F	2 3
267		AZ	PPD		4443	15	06/21/91		51
268		AZ	PPD	263	3839	7		27 M	51
269		ΑZ	PPD -	269		10			24
270		AZ	PPD	270			06/02/91		56
271		ΔZ	665	271		15			26
272		61	PPD	~ ~	3796	C 4 4	06/24/91 07/04/91		26 26
273		AZ	PPD PPD	272 273					2 4
274		AZ AZ	FFD CR3		63839		· · · · · · · · · · · · · · · · · · ·	•	26
275 274		AZ	PPD		4293	, 2			2 6
4/9 277			erj	276		e e e e e e e e e e e e e e e e e e e			등 김
		j ⊼≞	ere	-	0328	4	07/03/91		2 4
277			Fr.	273			08/21/53	,	2 5
د مید میچند معنو			270	277		2		医疗 网络	
201		et 2		220	8 4793 -	~	: cercaker		- ,
131	1:0223			281	. 459°	-			
122	\$ 100224	•	್ರಾ			-	. 23/27/21		
2.6-	تَشْتَنْهُمُ مِنْ اللَّهُ اللَّهُ ا	AI	623			14	00/27/91	1 49 5 	2.7
233		22	59D		, 1 393) <u>ng/75/19</u> 1 1 10 (10 (10 (10 (
	4 100237	ĄZ	rrd		5 670.		1 02/12/9:		3 5
	7 100230	AZ	PPD		5 3184		2 02/03/91		. .
28:	3 100287	A1	PFD	287	7 4037	-110	3 09/23/9:	1 32 M	2. J.

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DIRECTORY OF DATABASE 03/18/94

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PAGE NO. 03/18/94 7

RECORD NUMBER	CONTROL NUMBER		ABENCY	AGENCY COUNT	DRE OFFICER	DRE COUNT	ARREST DATE	AGE SEX	A 0 C R =E =K
		AZ	PPD	268	4545	2	09/20/91	65 M	2 1
	100290	-		289	4149	13	09/17/91	33 M	2 3
290	100291	AZ	PPD		4593		10/30/91	57 M	2 l
291	100272	67	PPD	291	3807	15	10/02/91	99 M	5 2
272	100292	AZ	opd 1			, ,	10/28/91	33 M	26
273	100274	ΑZ	PPD		3437	11	10/16/91	37 M	2 3
294	100275	A2	PPD		4293	3	10/13/91	42 M	2 6
295	100296	AZ	PPD	224	3701		10/13/91	23 M	26
296	100297	AZ	PPD		3398	20		22 M	2 6
297	100298	AZ	PPD	296		12	10/09/91		. 2 &
298	100299	AZ	:PD		4017	10	10/09/91		· 4 ~
299	100300	AZ	PPD	273		3	10/08/91		2 6
300	100301	AZ	:PPD	299	4337	1,1	10/04/91	37 M	4445364
301	100302	AZ	PPD		4593	5	10/03/91	28 F	2 9 5
	100303	AZ	PPD	301	4017	11	10/02/91	24 M	26
	100304	AZ	PPD	302	3304	11	10/04/91		2 6
	100305	AŻ	9PD	303	3664	2	11/08/91	99 M	2 5 2 6 2 6
305		AZ	29D	304			11/27/91		2 6
306		AZ	PPD	305	2534	22	11/30/91		26
	100308	AZ	PPD	306	4228	11	11/27/91		26
309		AZ	PD	307	4228	· 12			24
	100310	AZ	PPD	308	4228	13	11/20/91		2 6
	100311	AZ	PPD	309		10			26
		AZ	PPD		2835	13			26
311		AZ	PPD		3664	3			24
312		AZ	PPD		4228	14	11/05/91	. 28 M	56
	100314	AZ	PPD		4147	17	11/04/91	L 45 F	26
	100315	AZ	PPD		3839	10	12/31/91	(31 F	2 6
	5 100316	AZ	PPD		3839	11	12/26/91	L 21 M	3 i
316		AZ	PPD		4339	12	12/14/91	143 F	2 5
	7 100318	AZ	PPD		3437	11			26
	3 100319	AZ	PPD		3 3701	10			3 6
319			99D		3701	11	12/12/93	1 20 F	26
) 100321	AZ AZ	PPD		4017	12			22
	100322		PPD	321		12			26
32:		AZ AZ	PPD PPD	323	-	17			26
32:			rru Dad	323		5	12/05/93	1 33 M	21
	4 100325	AZ	PPD	27	4 3837	12	2 12/02/9		21
	5 100326	AZ			5 3125		09/25/90		26
	6 100327	AZ	PPD		6 3398		1 09/18/9		26
	7 100328	AZ			6 3378 7 4545		02/14/9		2 4
	8 100329	AZ	2PD				9 03/19/9		36
	9 100330	AŻ			3 4037 7 4228 '		5 06/13/9		Ŝ è
	0 100331	AZ			9 4228 0 4149		0 04/27/9		ŽŽ
	1 100332	AZ					3 02/01/8		2 6
	2 100333	AZ		33	1 2534		6 10/14/9		$\overline{2}$
	3 100334	AZ			2 4393		2 03/07/9		$\frac{1}{2}$ $\dot{\epsilon}$
	4 100335	AZ			3 4293		2 09/07/9		$\overline{2}$
33	5 100336	AZ			4 3701		2 03/11/9		2 è
33	6 100337	AZ	PPD	33	5 3398	£.		,	

DIRECTORY OF DATABASE 03/18/94

1

PAGE NO. 03/18/94 8

RECORD CONTROL NUMBER NUMBER	STATE AGENCY	AGENCY DRE COUNT OFFICER		A 0 AGE SEX C R === === =E =K	č
337 100338	AZ PPD	336 4192	12 10/25/90	65 F 2 6	5
338 100339	AZ PPD	337 4192	13 01/05/91	25 F 2-6 20 F 2-6	_
339 100340	AZ PPD	338 3701	13 06/25/91		
340 100341	AZ PPD	339 3525	6 09/06/90 17 04/10/91	22 M 2 6 22 M 2 6	
341 100342	AZ PPD	340 3807 341 3901	10 05/07/91	33 M 2 4	
342 100343	AZ PPD AZ PPD	342 4545	5 09/20/92	33 M 2 4	ł
343 100344	AZ PPD	344 4192	14 10/21/89	25 M 2 &	5
344 100346 343 100347	AZ PPD	345 3528	5 10/28/87		\$
345 100348	AZ PPD	346 3528	\$ 10/27/89		5
347 100349	AZ PPD	347 4443	18 10/23/37		5
348 100350	AZ PPD	348 3398	22 10/22/89		5 6
349 100351	AZ PPD	349 4443	19 10/20/89 20 10/19/89		5
350 100352	AZ PPD	350 4443	20 10/19/89 21 10/20/89		6
351 100353	AZ PPD	351 4443 352 3528	7 10/20/89		<u>6</u> .
352 100354	AZ PFD AZ PPD	352 3028	15 10/19/39	35 M 2 (6
53 100355 54 100355	AZ PPD AZ PPD	354 4228	16 10/12/39		6
354 100356 355 100357	AZ PPD	431 2705	5 10/13/89		6
356 100358	AZ PPD	356 3307	18 10/10/89		é,
357 100359	AZ PPD	357 229?	5 10/06/87		6
358 100360	AZ PPD	358 4228	17 10/06/89		6 6
359 100361	AZ PPD	359 4443	22 10/01/87		о А
360 100362	AZ PPD	360 2534	19 02/26/92		6
361 100363	AZ PPD	361 3807	13 02/27/92		6
362 100364	AZ PPD	362 3437 363 3901	11 02/28/92		4
363 100365	AZ PPD AZ PPD	363 3771	1 02/27/92	38 F 2	4
364 100366 365 100367	az PPD Az PPD	365 3807	20 02/25/92		ć
365 100368	AZ PPD	366 4149	21 02/21/92		6
367 100369	AZ PPD	367 3398	24 12/01/92	22 M 2	Ę
368 100370	AZ PPD	368 4037	30 02/09/92	50 M 2 32 F 2	1
369 100371	AZ PPD	367 3398	25 02/08/92 12 02/03/92	32 F 2 26 M 2	4 4
370 100372	AZ PPD	370 3901 371 4443	23 01/04/91	25 M 2	-
371 100373	AZ PPD	371 4443 372 3398	26 03/31/92	19 M 2	ė
372 100374	az PPD Az PPD	373 3385	11 03/30/91	36 F 2	ć
373 100375 374 100376	AZ PPD	374 3304	12 03/27/92		4
375 100377	AZ PPD	375 3304	13 03/27/72		ζ.
376 100378	AZ PPD	376 3125	7 03/24/92		ς.
377 100379	AZ PPD	377 2299	6 03/20/92		4
378 100380	AZ PPD	378 3996	8 03/18/92		4
379 100331	AŽ PPD	379 3701	14 03/18/92 14 03/10/92		4
380 1 003 82	A7 PPÖ	380 3304 381 3807	21 03/10/92		:
38: 100063	AZ PPD AJ PPD	381 4607 381 2 385	14 03/13/92	34 M 2	
382 100384 735 100383	AL PPD AZ FPD	352 3304	15 09/16/92	24 F 2	,
383 100393 384 100383	a: FPD	364 3304	16 03/09/92		:
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PAGE NO. 03/18/94

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17

DIRECTORY OF DATABASE 03/18/94

RECORD CONTRO NUMBER NUMBER	STATE AGENCY	AGENCY DRE COUNT OFFICER	DRE ARREST COUNT DATE	A O AGE SEX C R EEE EE EK
385 100387 386 100388	AZ PPD AZ PPD	385 3184 386 4293	3 03/08/92 13 03/03/92	33 M 2.6
387 100389	AZ PPD	387 3437	13 03/03/92	28 M 3 4 42 F 2 5
388 100390	AZ PPD	388 4192	17 04/30/92	
389 100391 390 100392	AZ PPD	389 4545	6 04/23/92	31 F 2 6 25 M 5 6
371 100372	AZ PPD	370 4149	22 04/17/92	40 M 2 6
372 100374	AZ PPD AZ PPD	391 3701	15 04/16/92	41 M 2 6
393 100395	AZ PPD	392 4593	7 04/09/92	25 M 3 6
394 100396	AZ PPD	393 3437 394 4192	15 04/10/92	64 M 2 6
395 100397	AZ PPD	375 3398	18 04/07/92	25 F 2 4
396 100393	AZ PPD	376 3125	27 04/06/92	22 F 2 6
397 100399	AZ PPD	397 4845	8 04/01/92 7 04/03/92	43 M 5 6
398 100400	AZ PPD	378 2885	15 03/31/92	21 M 5 4
379 100401	AZ PPD	377 3701	13 05/22/92	27 F 2 4 21 M 2 7
400 100402	AZ PPD	400 3701	16 05/19/92	
401 100403	AZ PPD	401 2299	7 05/16/92	
402 100404 403 100405	AZ PPD	402 3385	12 05/14/92	
404 100405	AZ PPD	403 4037	31 05/17/92	34 F 2 6 37 F 2 6
405 100407	AZ PPD	404 4545	3 05/14/92	25 M 2 6
406 100408	AZ PPD AZ PPD	405 3398	28 05/10/92	38 M 2 6
407 100409	AZ PPD	406 4293	14 05/05/92	$21 M = 5 \tilde{2}$
408 100410	AZ PPD	407 3996 408 4228	9 06/30/92	36 F 2 6
409 100411	AZ PPD	409 2885	18 06/20/92	32 M 2 6
410 100412	AZ PPD	410 2385	16 06/20/92	44 F 2 4
411 100413	AZ PPD	411 4443	17 06/19/92 24 06/17/92	42 M 2 3
412 100414	AZ PPD	412 3125	24 06/17/92 9 06/15/92	20 M 2 6
413 100415	AZ PPD	413 3525	7 07/22/92	35 M 2 6
414 100416	AZ PPD	414 3839	13 07/17/92	33 M 2 7 26 M 2 A
415 100417	AZ PPD	415 3807	22 07/17/92	26 M 2 6 37 M 2 1
416 100418 417 100419	AZ PPD	416 2885	18 07/03/92	37 F 2 4
413 100420	AZ PPD	417 3304	17 08/28/92	46 M 2 6
419 100421	AZ PPD AZ PPD	418 4545	9 08/22/92	36 M 2 6
420 100422	AZ PPD	419 3807	23 08/21/92	43 M 2 6
421 100423	AZ PPD	420 3839 421 3839	14 08/14/92	19 M 5 2
422 100424	AZ PPD	422 4293	15 08/08/92	23 M 5 6
423 100425	AZ PPD	423 3437	15 08/04/92 16 09/25/92	28 M 2 1 29 M 2 6
424 100426	AZ PPD	424 3837		29 M 2 6
425 100427	AZ PPD	425 4667	1 10/17/92	32 M 3 3 43 M 2 6
426 100428	AZ PPD	426 4667		
427 100429 428 100430	AZ PPD	427 3125		27 M 2 6 99 F 2 6
428 100430 429 100431	AZ PPD	428 4339		38M 2.6
430 100432	AZ PPD AZ PPD	429 3437	17 12/20/92	41 M 2 6
431 100433	az ppd Az ppd	432 4037	32 03/20/90	29 M 2 6
432 100434	AZ PPD	433 4149	23 08/21/91	33 M 2 1
	76 (FFU	434 3839	17 01/17/92	20 M 2 6

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PAGE ND. 10 03/18/94

DIRECTORY OF DATABASE 03/18/94

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	RECORD NUMBER	CONTROL NUMBER		AGENCY		DRE OFFICER	DRE COUNT	ARREST DATE	AGE	SEX	A C =E	O R ≓K	- <u>.</u>
2-9-9 ³⁻⁹ -9													
	422	100435	AZ	PPD	435	3839	13	08/24/91	41	м	2	6	
		100436	AZ	PPD		3730		07/21/91	52	M	2	1	
		100437	AZ	PPD		3528		06/06/91	39	м	2	ô,	
		100438	AZ	PPD		4228		05/02/90	37	м	2	6	
		100439	AZ	PPD		4443		07/27/91	42	М	2	5	
		100440	AZ	PPD		2885	19	07/27/91	42	М	2	i	
		100441	AZ	PPD	441	4293		12/06/91	25	M	2	4	
	440	100442	AZ	P P D	442	4228		11/22/91	- 29		2	4	
	441	100443	AZ	PPD		4667		01/07/93		M	2	4	
		100444	AZ	PPD		4192		01/07/93		F	2	1	
		100445	AZ	PPD		4228		01/12/93		M	2	3	
		100446	AŻ	PPD		3398		01/24/93	23	M	22	6 6	
		100447	AZ	PPD		4443		01/21/73	34 43	F M	2	6	
		100448	AZ	PPD	· · -	4545		01/30/93		M	2	4	
		100449	AZ	PPD PPD		4192 4667		01/09/93	28	M	2	Æ	
		100450	AZ	ppd Ppd		3125				м	ž	6	
		100451 100452	AZ AZ	PPD		3839		01/12/92		M	5	4	
		100453	AZ	PPD		4443		01/30/93			- 2	1	
		100454	AZ	PPD		4228		01/26/93		í T	5	6	
		100455	AZ	PPD		4228		01/12/93		М	2	1	
		100456	AZ	PPD		3125	12	02/18/93	34	M	2	\Leftrightarrow	
		100457	AZ	PPD		4443	28	02/01/93	20	M	2	7	
		100458	AZ	PPD	458	4545		02/12/93		М	2	4	
	457	100459	AZ	PPD		4192		02/09/93		М	- 2	1	
	453	100460	AZ	PPD		3125		02/05/93		М	2	6	
		100461	AZ	PPD		433?		02/05/93			2	6	
		100462	AZ	PPD		3304		02/05/93			2	6	
		100463	AZ	PPD		3398		02/07/93		M M	2 N (3	6	
		100464	AZ	PPB PPD		3125		02/23/93 02/27/93			3	6 4	
		100465	AZ	220 222		3839		02/25/93			2	е 6	
		100466	AZ	PPD PPD		3996 3385		02/25/93			$\frac{4}{2}$	- 1	
		100467 100468	AZ AZ	PPD		3839	21				2	4	
		100469	AZ	PPD		4545		03/29/93			2	1	
		100489	AZ	PPD		4443	29				2	ŝ	
		100471	AZ	PPD		3996		03/31/93			2	6	
		100472	AZ	PPD		4593		03/12/93		М	2	6	
		100473	AZ	PPD		3398		03/07/93		F	- 2	- 6	,
	472	100474	AZ	PPD	474	4667	5	03/09/93			2	6	
	473	100475	AZ	PPD		4593	9				- 2	- 6	,
		100476	AZ	PPD		4593		03/16/93			2	1	
		100477	AZ	FPD		3664	-	04/15/93	_		- 2		
		100478	AZ	PPD		3125		04/09/93			2	- 6	
		100479	AZ	PPD		4575		04/28/93			5		
		100480	AZ	PPD SSD		4293		04/09/93			3		
		100481	AZ	PPD		3807		04/16/93			22		
	480	100482	A7	PPD	452	3398	25	04/24/93	· -21	r.,	4	6	

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PAGE NO. 03/18/94

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11

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DIRECTORY OF DATABASE 03/18/94

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RECORD NUMBER	CONTROL NUMBER	STATE	AGENCY	AGENCY COUNT	DRE OFFICER	DRE COUNT	ARREST DATE	AGE 	SEX	A C =£	0 R ≓K
481	100483	AZ	PPD	483	4593	11	04/09/93	30	F	2	6
432	100484	AZ	PPD	434	3839	22	04/20/93	32	М	2	4
433	100485	AZ	PPD	485	3437	18	04/22/93	43	М	2	6
484	100486	AŻ	PPÖ	436	4593	12	04/03/93	39	M	3	- 6
485	100487	AZ	PPD	437	4545	13	04/16/93	29	М	5	4
436	100489	AZ	₽₽D	489	3579	3	05/05/93	38	F	2	8
437	100470	AZ	FPD	490	4339	15	05/21/93	23	M	5	4
428	100491	AZ	PPD	491	4593	13	05/01/93	62	21	22	1
489	100492	AZ	PPD	492	4667	- 6	05/18/93	57	М		6
490	100493	AZ	PPD	493	3664 .	5	05/17/93	46	м	2,	1
491	100494	AZ	PPD	494	4667	7	05/20/93	45	Μ	2	-6
492	100495	A2	PPD	495	3839	23	05/24/93	25	М	2	7
473	100496	AZ	PPD	496	3996	12	02/08/93	41	М	2	6
494	100497	AZ	PPD	497	3437	19	05/23/93	29	F	2	6
495	100498	AZ	PPD	498	3398	33	07/11/92	- 27	М	2	6
496	100499	AZ	ppd	499	3125	17	07/17/91	- 22	М	5	6
497	100501	AŻ	PPD	500	2885	20	10/12/91	34	f	2	6
498	100502	AZ	PPD	501	4293 -	18	06/12/92	23	F	2.	6
499	100503	AZ	PPD	502	3385	14	05/15/92	45	M	2	1
500	100504	AZ -	PPD	503	3125	18	03/09/93	42	M	2	6

APPENDIX V

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"OTHER" DRUGS REPORTED

FROM

ANALYSIS OF SPECIMENS

"OTHER" DRUGS: DRUGS DETECTED IN URINE AND BLOOD SPECIMENS FOR WHICH THERE ARE NO CHECKBOXES

	Classificati	ion							
Drug or Metabolite	<u>(See Key</u>)							
AZACYLONOL									
BENZTROPINE									
AMITRIPTYLINE	P								
CARBAMAZEPINE	Р								
CARISOPRODOL	P								
CHLORPHENIRAMINE	P								
CLOMIPRAMINE	-								
DESIPRAMINE	P+M	See Note 1							
DIPHENHYDRAMINE	Р	000.1012 .							
DOXYLAMINE	P								
DOXEPIN	P								
DESMETHYLDOXEPIN	Ň								
EPHEDRINE	P								
FLUOXETINE	P								
HYDROCODONE	P								
3-HYDROXY-N-METHYLMORPHINAN	M								
LIDOCAINE	P								
MEPROBAMATE	P+M	See Note 2							
METHADONE	P	000 11000 2							
METHORPHAN	P								
MEPERIDINE	P								
NOREPHEDRINE	M								
NORPSUEDOEPHEDRINE	M								
NORPROPOXYPHENE	M								
NORCHLORPHENIRAMINE	M								
NORCODEINE	M								
PSUEDOEPHEDRINE	P								
PROPOXYPHENE	P								
NORTRIPTYLINE	P+M	See Note 3							
PROMETHAZINE	P								
OXYCODONE	P								
PRIMIDONE	P	-							
6-MONOACETYLMORPHINE	M								
TEGRETOL	P								
TRAZODONE	P								
TOLUENE	P								
	•								
Classifi	ication Key:								
Parent	Drug = P								
Metal	polite = M								
May be either $= P + M$									

May be either = P + M

Note:

If imipramine is present, desipramine is a metabolite.
 If carisoprodol is present, meprobamate is a metabolite.
 If amitriptyline is present, nortriptyline is a metabolite.