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LIST OF SUPPLIERS OF CHEMICALS AND EQUIPMENT

Acrylamide	BDH, UK
ACD vacutainer tubes	Becton Dickinson, UK
Agarose	FMC Bioproducts, USA
Ammonium Chloride	Sigma, USA
Ammonium Persulphate	Stratagene, USA
B-mercaptoethanol	Merck, Germany
Beckman DU®-65 spectrophotometer	Beckman, USA
Biorad Model 200/2.0 Power Supply	Biorad, USA
Bisacrylamide	BDH, UK
Blue/orange loading dye	Promega, USA
Boric Acid	USB, USA
Bromophenol blue	BDH, UK
BRL model S2 electrophoresis apparatus	BRL, USA
Bovine Serum Albumin	Roche, Germany
4-chloro-1-naphthol (C ₁₀ H ₇ ClO)	Sigma Chemical Company, USA
Conical polycarbonate centrifuge tubes	Nunc, Denmark
Coomassie brilliant blue R-250	BDH, UK
Coomassie Plus Protein Assay Reagent Kit	Pierce, USA
100bp DNA ladder	Promega, USA
EDTA	Roche, Germany
Ethanol	BDH, UK
Ethidium Bromide	Roche, Germany
Eppendorf tubes	Eppendorf, Germany
Extran	Merck, Germany
EZCast Gel Casting Boot	BRL, USA
Filter tips	QSP, USA?
Filter paper	Whatman, USA

Gel 3-place casting trough	CBS Scientific, USA
Glacial acetic acid	Saarchem, RSA
Goat anti-rabbit antibody	Roche, Germany
GS300 Transmittance/Reflectance Scanning Densitometer	Hoefer Scientific Instruments, USA
Hägar Designs HM2 centrifuge	Hägar Designs, RSA
Heating block	Hägar Designs, RSA
Hoefer GS300 Transmittance/Reflectance Scanning Densitometer	Hoefer Scientific Instruments, USA
Hoefer Red Rotor Model #PR70-230V	Hoefer Scientific Instruments, USA
Hoefer SG Series Gradient Maker	Hoefer Scientific Instruments, USA
Hoefer Sturdiel Vertical Slab Gel Unit (SE400)	Hoefer Scientific Instruments, USA
Horizontal mini-gel kit, model #MGU-200T	CBS Scientific Company, USA
Hybond-C Super Transfer Membranes	Amersham, USA
Hydrochloric Acid (HCl)	Saarchem, RSA
Hydrogen Peroxide (H ₂ O ₂)	Saarchem, RSA
Isopropanol	Saarchem, RSA
Jouan BR3.11 centrifuge	Jouan, France
MDE gel matrix	FMC Bioproducts, ISA
Methanol	Saarchem, RSA
Mineral oil	Sigma, USA
Na ₂ HPO ₄ ·2H ₂ O	Saarchem, RSA
NaCl	Saarchem, RSA
NaH ₂ PO ₄	Saarchem, RSA
NaOH	Saarchem, RSA
Oligonucleotide primers	IDT, USA Inqaba Biotec, RSA
Parafilm	Whatman, USA
PCR Core Kit	Roche, Germany

PCR Master Kit	Roche, Germany
PCR Master Mix	Promega, USA
PCR tubes	QSP, USA?
Perkin Elmer Cetus DNA Thermal Cycler	Perkin Elmer, USA
[α - ³² P]dATP	Amersham Biosciences, UK
PS500X DC power supply	Hoefer Scientific Instruments, USA
SDS	BDH, UK
Sequenase version 2.0 DNA Sequencing Kit	Amersham Biosciences, UK
Sorvall RMC-14 centrifuge	Sorvall, USA
Stop Solution	Promega, USA
TEMED	Promega, USA
Tris	BDH, UK
Urea	GibcoBRL, USA
UV transilluminator	UVP, USA
Whatman #3 Filter Paper	Whatman, USA
X-ray film	Agfa, Germany