



Chemotherapy in Non-Small Cell Lung Cancer: An Update

A meta-analysis of randomised trials using individual patient data

Protocol E

Comparison 7

Supportive care vs supportive care + chemotherapy

Conducted by the Non-Small Cell Lung Cancer Collaborative Group (NSCLCCG)

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Introduction

More than a million new cases of lung cancer are diagnosed each year¹. About 80% of these tumours are of non-small cell histological type², including adenocarcinomas, squamous cell and large cell carcinomas. Non-small cell lung cancer (NSCLC) is the main cause of deaths from cancer and five-year survival across all stages of disease is about 14%³.

Surgery is generally regarded as the best treatment option, but only about 30% of tumours are suitable for potentially curative resection⁴. A further 20% of patients, usually those presenting with locally advanced disease, undergo radical thoracic radiotherapy. The remaining 50% of patients, with late stage or metastatic disease, are usually treated palliatively.

In 1991, the British Medical Research Council's Cancer Trials Office (MRC), Cambridge; the Institut Gustave Roussy (IGR), Villejuif, France; and the Istituto Mario Negri (IMN), Milan, Italy initiated an individual patient data (IPD) meta-analysis to assess the role of chemotherapy in the treatment of NSCLC. At that time, despite over thirty years of research involving more than 9000 patients in over 50 randomised clinical trials (RCTs), the efficacy of chemotherapy, when combined with local treatment or supportive care, was unclear. With few exceptions, most trials were too small to reliably detect moderate treatment effects. Consequently, although a few trials reported significant results, both for and against chemotherapy, most trials were inconclusive.

This IPD approach to meta-analysis and systematic review involved the central collection, validation and analysis of the original trial data. It did not rely on data extracted from publications. At the outset, the secretariat contacted the investigators responsible for each trial and established the NSCLC Collaborative Group on whose behalf the meta-analysis was carried out and published in the British Medical Journal in 1995⁵. This has become a 'landmark publication' and was cited nearly 900 times between 1995-2003.

The meta-analysis concluded that despite previous scepticism and controversy, modern chemotherapy could have a role in treating NSCLC. In particular, there was strong evidence that for more advanced disease, chemotherapy given in addition to radical radiotherapy and given in addition to best supportive care, prolonged survival. The results for early stage disease, although in favour of chemotherapy, were less clear-cut. A fuller presentation of the results of the meta-analysis published in 1995 is given in Box 1.

Modern cisplatin-based regimens may offer the first effective adjuvant treatment in NSCLC and this should be evaluated in a prospective large-scale trial. New agents may offer further advantages and should be explored.

Since the meta-analysis was published, there has been renewed enthusiasm for investigations of chemotherapy in NSCLC and a considerable number of new RCTs have been completed. The total number of patients randomised has risen from 9387 to around 23000 patients. In particular, there have been many new trials in the surgical setting including trials of neoadjuvant chemotherapy. A number of new agents and timings have been investigated in all settings. As the aim of the NSCLC Collaborative Group is to provide an up to date and reliable review of the role of chemotherapy, both to act as a sound basis for evidence based medicine and to help guide future research, it is now timely to undertake a major update and re-evaluation of the 1995 meta-analysis.

Seven therapeutic comparisons will be explored:

- 1 surgery vs surgery + adjuvant chemotherapy
- 2 surgery vs neoadjuvant chemotherapy + surgery
- 3 surgery + radiotherapy vs surgery + radiotherapy + adjuvant chemotherapy
- 4 radiotherapy vs radiotherapy + sequential chemotherapy (neoadjuvant before radiotherapy- and/or adjuvant –after radiotherapy) or alternated radio-chemotherapy
- 5 radiotherapy vs radiotherapy + concomitant chemotherapy
- 6 radiotherapy + sequential chemotherapy vs radiotherapy + concomitant chemotherapy
- 7 supportive care vs supportive care + chemotherapy

Three of these comparisons (2, 5 and 6) are new; this reflects changes in practice and interest since the 1995 Meta-analysis and ensures that this systematic review is as inclusive and comprehensive as possible.

For clarity a separate protocol has been produced for each individual comparison, each of which can be considered as an independent meta-analysis, but when considered together will allow us to evaluate the overall picture of chemotherapy in non-small cell lung cancer.

This protocol relates to **comparison 7** of the meta-analysis, copies of all protocols are available on request or can be downloaded from http://www.ctu.mrc.ac.uk/download.asp or http://www.igr.fr/php/index.php?ids_path=2.51.70.127.567

The meta-analyses will be jointly run by the Medical Research Council (MRC) and the Institut Gustave-Roussy (IGR). See Appendix D for further meta-analyses completed by these two groups.

Box 1

Summary of Results of 1995 Meta-analysis¹

The main objective of the meta-analysis was to investigate the effect of chemotherapy on survival when given in addition to appropriate local treatment:

Early disease

surgery versus surgery + chemotherapy

surgery + radiotherapy versus surgery + radiotherapy + sequential chemotherapy *Locally advanced disease*

radical radiotherapy versus radical radiotherapy + chemotherapy

Advanced disease

supportive care versus supportive care + chemotherapy

Trials were classified as belonging to one of four pre-specified categories of chemotherapy -Regimens containing cisplatin

-Regimens using long-term alkylating agents (but not cisplatin)

-Regimens containing etoposide or vinca alkaloids (but not cisplatin)

-Other regimens

Results

Comparison	Hazard Ratio	Confidence Interval	p-value
Surgery vs surgery + chemotherapy			
Long-term alkylating agents	1.15	1.04-1.27	0.005
Other drugs	0.89	0.72-1.11	0.30
Cisplatin based	0.87	0.74-1.02	0.08
Surgery + RT vs surgery + RT + chemotherapy			
Long-term alkylating agents	1.35	0.83-2.20	0.23
Cisplatin based	0.94	0.79-1.11	0.46
Radical RT vs radical RT + chemotherapy			
Long-term alkylating agents	0.98	0.83-1.16	0.81
Vinca-alkaloids	0.87	0.70-1.09	0.23
Other drugs	0.98	0.74-1.29	0.88
Cisplatin based	0.87	0.79-0.96	0.005
Supportive care vs supportive care + chemotherapy			
Long-term alkylating agents	1.26	0.96-1.66	0.095
Vinca-alkaloids / etoposide	0.87	0.64-1.20	0.40
Cisplatin based	0.73	0.63-0.85	<0.0001

A further objective was to assess whether any possible effects were consistent in the subgroups of age, sex, extent of disease, tumour stage, histology and performance status. Subgroup analysis of trials using cisplatin based regimens found no indication that any particular type of patient benefited more or less from chemotherapy.

Conclusions

The results were consistent across primary treatment settings and they tended to show a benefit of modern cisplatin-based chemotherapy regimens although essential drugs were not identified.

¹ Non-small Cell Lung Cancer Collaborative Group. Chemotherapy in non-small cell lung cancer: a meta-analysis using updated data on individual patients from 52 randomised clinical trials. British Medical Journal 1995;311:899-909.

Methods

Design

A systematic review and quantitative meta-analysis based on updated individual patient data will be carried out. This approach involves the central collection, validation and analysis of data from all patients from all relevant randomised trials.

Treatment comparisons

Seven meta-analyses will be carried out over three main clinical settings, corresponding to the stage of disease and to the primary treatment

Surgery

- 1 surgery vs surgery + adjuvant chemotherapy
- 2 surgery vs neoadjuvant chemotherapy + surgery
- 3 surgery + radiotherapy vs surgery + radiotherapy + adjuvant chemotherapy

Radiotherapy

- 4 radiotherapy vs sequential radiotherapy + sequential chemotherapy
- 5 radiotherapy vs radiotherapy + concomitant chemotherapy
- 6 radiotherapy + sequential chemotherapy vs radiotherapy + concomitant chemotherapy

Supportive care

7 supportive care vs supportive care + chemotherapy

Identification of trials

There is good evidence that investigators and journals alike are more likely to publish trials with positive results⁶⁻⁸. In order to avoid such publication bias, both published and unpublished trials will be included in the meta-analysis. To identify as many relevant trials as possible, systematic searches of a number of trial sources will be carried out and updated during the course of the project, ensuring a comprehensive and up-to-date database of trials.

Electronic Databases

The optimum search strategy for retrieving randomised controlled trials (RCTs) from Medline, developed by the Cochrane Collaboration⁹ will be modified (Appendix B).

- To specifically retrieve RCTs of chemotherapy for NSCLC
- And used to search Medline and Cancerlit

In addition the following electronic bibliographic databases will be searched.

- The Cochrane Central Register of Controlled Trials (CENTRAL)
- Proceedings of ASCO 1995 2003

Trial Registers

Trial registers will be used to supplement searches of electronic databases with trials that may not (yet) be published or are still recruiting patients:

- UKCCCR Trials Register
- ClinicalTrials.gov
- Physicians Data Query Protocols (open and closed)
- Current Controlled Trials 'metaRegister' of controlled trials

Hand Searches

The following hand searches will be carried out with the aim of identifying trials that may have only been reported as abstracts or that might have been missed in the searches described above:

- Proceedings of the American Society for Clinical Oncology (ASCO) 1993-1994
- Proceedings of the IASLC World Lung Cancer Conference 1997-2003
- Proceedings of the European Society of Medical Oncology (ESMO) 1996 2002
- Proceedings of the European Cancer Conference Organization (ECCO) 1995 2003
- Bibliographies of all identified trials and review articles will be searched

Experts in the field

All participating trialists will be asked to review and supplement a provisional list of trials

Eligibility criteria

7 supportive care vs supportive care + chemotherapy

Trials must

- be properly randomised in a way which precludes prior knowledge of treatment assigned
- be unconfounded
- have commenced randomisation on or after January 1st 1965
- have completed accrual (ongoing trials are listed but no data collected) before 2003
- have included patients with non-small cell lung cancer

Patients must

- be unsuitable for surgery
- be unsuitable for radical radiotherapy
- be randomised to receive supportive care and chemotherapy or supportive care alone
- not have received prior chemotherapy
- not have had any prior malignancy

Data Collection

New Trials

For all new trials, basic survival and baseline characteristics will be sought for **all** patients randomised into each trial. Up to date follow-up will be requested in order to report on both short and longer-term outcomes.

patient identifier (preferably not patient name) date of birth or age at randomisation sex performance status tumour stage (TNM) histology date of randomisation treatment allocated (specify each arm of chemotherapy if several regimens used) survival status date of last follow-up date of death cause of death whether excluded from trial analysis reason for exclusion

Trials already included in the 1995 Meta-analysis

Trials of long-term alkylating agents will not be updated as these are old trials that provided mature data for the 1995 analysis.

Most of the trials included in the 1995 meta-analysis provided mature data and we do not anticipate that much additional information will be available. However, if trialists are able to provide any additional updated information, this will be included in the new analyses.

Suggested coding conventions for these data are provided (Appendix C) and although using them will facilitate data transfer, it is not essential. Data will be accepted in whatever format is most convenient for the individual trial investigator or data centre and can be supplied by email, computer disk, on data collection forms or as a computer printout. We will also ask for a limited amount of information on trial design as well as the original trial protocol, associated on-study forms and publications

A final copy of the data from each trial will be returned to the trialists for verification. The data collation and checking for **this comparison** will be done by the **MRC**. Copies of the final agreed database of all trials included in all comparisons will be held by the MRC and the IGR. All trial data will be held securely and will not be used, circulated or distributed in any way that allows access to individual trial data, without first seeking permission from trial investigators.

Analysis

The ultimate aim will be to obtain and analyse data from all randomised patients included in all of the relevant randomised trials.

The analysis will be performed on the endpoint of overall survival. All analyses will be of randomised patients and will be carried out by intention to treat, that is, patients will be analysed according to the treatment allocated, **irrespective** of whether they received that treatment. Survival analyses will be stratified by trial, and the log-rank expected number of deaths and variance will be used to calculate individual and overall pooled hazard ratios by the fixed-effect model¹⁰. Thus, the times to death for individual patients will be used within trials to calculate the hazard ratio, representing the overall risk of death for patients who were allocated chemotherapy compared to those who were not.

Results will also be presented as absolute differences at relevant time points calculated from the hazard ratio and baseline event rate for patients not allocated chemotherapy;¹¹ proportional hazards are assumed. Confidence intervals for absolute differences will be similarly calculated from the baseline event rate and the hazard ratio at the 95% CI boundary values. χ^2 heterogeneity tests⁵ will be used to test for gross statistical heterogeneity, the I² statistic¹² will be used as a measure of consistency. Survival curves will be presented as simple (non-stratified) Kaplan-Meier curves¹³. All p-values will be two-sided.

Analyses by trial level characteristics

The effect of chemotherapy may vary across trials in the meta-analysis because they have each applied treatment in different ways. To explore this further, providing that there are sufficient data available, analyses are planned whereby trials, or arms within trials, will be grouped according to the type of chemotherapy regimen to determine whether there are any differences in treatment effect between these groups.

Trial characteristics will be reported in tabular form, information will include patient numbers, period of recruitment, treatment details and proportion of patients who received second-line treatment.

Type of chemotherapy

It is not practical to look at groups of trials using only exactly the same regimens, nor is it appropriate or sensible to look overall at all trials. We therefore plan to split trials into broad groupings according to the type of chemotherapy used. This will build on the groupings used in the 1995 meta-analysis, which revealed that old trials using long-term oral alkylating agents had a detrimental effect whereas trials using modern regimens showed a beneficial effect

Within each main treatment comparison, trials will be grouped by the type of chemotherapy regimen. If there are insufficient numbers of patients within any categories, categories may be combined.

Platinum based regimens

platinum + vinca alkaloid / etoposide platinum + anti-metabolic agent (e.g. tegafur, uft) platinum + taxane other platinum regimen

Non-platinum based regimens

vinca alkaloid / etoposide only anti-metabolic agent only taxane only other non-platinum regimen

Long-term alkylating agents

Owing to the results of the 1995 meta-analysis, long-term alkylating agents will be considered separately from other trials. Owing to their antiquity, these will not be updated and results will remain unchanged but will be included for completeness.

For these analyses a hazard ratio will be calculated for each trial and a pooled hazard ratio calculated for each treatment category. A test for interaction will be used to investigate if there are any substantial differences in the effect of treatment between these treatment categories. If there is no clear evidence of heterogeneity results may also be combined over categories.

Sensitivity Analyses

Hazard ratios for overall survival will also be calculated using a random effects model.

Hazard ratios for overall survival will also be calculated excluding any trials that are clear outliers.

Analyses by patient level characteristics

Providing there are sufficient data available, we will investigate whether any observed treatment effect is consistent across well-defined patient subgroups. These analyses will be carried out on all trials (except trials of long-term alkylating agents) and will be stratified by trial. If there are substantial heterogeneity and differences of effect between treatment categories, then subgroup analyses will be done within treatment categories.

If there are insufficient numbers of patients within any patient categories, categories will be combined. Chi-squared tests for interaction or trend will be used to test whether there is any evidence that particular types of patients benefit more or less from chemotherapy.

The subgroups are as follows:

Age (<60, 60-64, 65-69, 70+) Sex (Male, Female) Performance Status *(Good, Poor) Histology (Adenocarcinoma, Squamous, Large cell, Other) Stage **See below for calculation

*Performance Status

Meta-analysis Performance Status	WHO / ECOG	Karnofsky
Good	0, 1	100, 90, 80, 70
Poor	2, 3, 4	60 - 10

**Stage

Meta-analysis Stage / ISS 1986	(p)TNM Classification			AJCC Stage	UICC stage 1997
	(p)T	(p)N	(p)M		
I	0,1,2,X,S	0	0		IA, IB
II	0,1,2,X,S	1	0	II	IIA, IIB without T3N0
IIIA	a) 3	a) 0-1	0	III non metastatic	IIIA + T3N0
	b) 1-3	b) 2			
IIIB	4, Any N	3, Any T	0	III non metastatic	IIIB
IV	Any	Any	1	Any metastatic	IV

Alternative exploratory analysis

Given the large number of trial and patient characteristics of interest, there may be interactions between them that could potentially confound these analyses. If we encounter substantial heterogeneity within any of the seven main meta-analyses we will further explore the potential influence of these factors using multi-level modelling techniques.

This modelling aspect of the project will be developed in collaboration with Dr Julian Higgins from the MRC Biostatistics Unit, who is a member of the International Advisory Group.

Publication Policy

The results of the meta-analyses will be published and presented in the name of the NSCLC Collaborative Group comprising trialists contributing data for analysis, the Secretariat and Advisory Group. Following publication in a peer reviewed journal, the meta-analyses will be submitted to the Cochrane Library to appear in the Cochrane Database of Systematic Reviews.

The seven meta-analyses will be analysed separately. We aim to have one collaborators meeting, at which the results will be presented, but the comparisons may be published separately.

The IGR carried out an individual patient data meta-analysis of concomitant chemotherapy (cisplatin or carboplatin) and radiotherapy in locally advanced non-small cell lung cancer. The results were presented at the 2003 World Conference on Lung Cancer in the name of the MAC3-LC (meta-analysis of cisplatin/carboplatin based concomitant chemotherapy in non-small cell lung cancer) Group¹⁴. This study will be published in the name of this group. The update of this meta-analysis as well as the updating of the comparison on sequential radio-chemotherapy versus radiotherapy alone (comparison 4) and the direct comparison of these two types of radio-chemotherapy (comparison 6) will be published in the name of the current collaborative group.

Project Administration

As for the 1995 Meta-analysis, the MRC and the IGR will share project administration.

Comparisons 1, 2 & 7

The MRC will be responsible for all contact, data collection, verification and analysis for these comparisons.

Comparisons 3, 4, 5 & 6

The IGR will be responsible for all contact, data collection, verification and analysis for these comparisons.

Contacting Trialists

New trials

Trialists will be contacted, informed of the project, invited to collaborate and asked to supply data as outlined in the methods section.

Trials already included in the NSCLCCG Database

Trials of long-term alkylating agents will not be updated as these are old trials that provided mature data for the 1995 analysis.

Most of the trials included in the 1995 meta-analysis provided mature data and we do not anticipate that much additional information will be available. However, if trialists are able to provide any additional updated information, this will be included in the new analyses.

Timetable

Spring 2004	Write to Trialists seeking collaboration
Spring 2004-Spring 2005	Collate, check and verify incoming data, analyse individual trials
Summer 2005	Analyses
Autumn 2005	Present results to Trialists

Comparison 7

supportive care vs supportive care + chemotherapy

Trial	Period of recruitment	Drugs used	Number of patients
Long Term Alkylating Age	nt		
Oxford ¹⁵	1970-73	Pc, NM, Vb, Pn	188
Quebec ¹⁶	1978-79	Mx, A, Cy, C	38
		Subtotal	220
Platinum + vinca alkaloid /	etoposide		
NCIC CTG BR5 ¹⁷	1983-86	C, Vd, C, A, Cy	150
RLW 8351 ¹⁸	1982-86	C,Vd	167
Southampton ¹⁸	1983-86	C, Vb	32
NRH ¹⁹	1983-87	C, Et	87
UCLA ²⁰	1984-86	C, Vd	63
Ancona 1 ²¹	1985-88	C, Cy, E, Mi, Et, C	128
Cep-85 ²²	1985-88	C, Vd	49
*CRC TU LU3002 (MIC 2)23	1988-96	Mi, If, C	351
*Thongprasert et al ²⁴		(If, E, C) or (C, Mi, Vb)	287
*BLT ²⁵	1995-01	(C, Vd) or (C, Nv) or (C, Mi, If) or (C, Mi, Nv)	725
*Ancona 2 ²⁶		C, Et	47
*Helsing et al ²⁷	1990-95	Cy, Et	49
<u> </u>		Subtotal	1968
Other platinum regimens			
Other platinum regimens AOI-Udine ²⁸	1984-86	C, Cy, Mi	102
		Subtotal	102
Vinca-alkaloid / etoposide	only		
Gwent 2 ²⁹	1982-84	Et	186
*ELVIS ³⁰		Nv	191
		Subtotal	377
Anti-metabolic agent only			
Anti-metabolic agent only *Anderson et al ³¹	1994-96	G	300
	1	Subtotal	300
Taxane only			
*Ranson et al ³²	1995-97	Px	157
*Roszkowski et al ³³	1995-(98?)	Dx	207
	/ /	Subtotal	364
		Total	3331

* Trials completed since 1995 Meta-analysis

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Appendix A

Glossary of Drugs

A B Bu C Cb Cy Dx E Et G If Lo Me Mi Mx MXa NM OF Pc Pe Pm Pr Px Tg Ti Tn	Doxorubicin Bleomycin Busulphan Cisplatin Carboplatin Cyclophosphamide Docetaxel Epirubicin Etoposide Gemcitabine Ifosfamide Lomustine / CCNU Mesna Mitomycin C Methotrexate Mitoxantrone Nitrogen Mustard Oral Ftorafur Procarbazine Pepleomycin Porfiromycin Prednisolone Paclitaxel Tegafur Tirapazamine Teniposide
	•
••	
U	UFT (Tegafur + uracil)
Vb	Vinblastine
Vc	Vincristine
Vd	Vindesine
Vn	Vinorelbine

Appendix B

Search Strategy for Medline

- 1 Randomized Controlled Trial.pt.
- 2 exp Randomised Controlled Trials/
- 3 exp Random Allocation/
- 4 exp Double-Blind Method/
- 5 exp Single-Blind Method/
- 6 1 or 2 or 3 or 4 or 5
- 7 clinical trial.pt.
- 8 exp Clinical Trials/
- 9 clin\$ with trial.ab,ti.
- 10 (sing\$ or doubl\$ or trebl\$ or tripl\$ with blind\$ or mask\$).ab, ti.
- 11 exp Placebos/
- 12 placebo\$.ab.ti.
- 13 random\$.ab,ti.
- 14 exp Research Design/
- 15 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14
- 16 exp Carcinoma/
- 17 exp Lung Neoplasms/
- 18 (lung adj carcinoma\$).ab.ti.
- 19 (lung adj cancer\$).ab,ti.
- 20 (lung adj neoplasm\$).ab,ti.
- 21 cancer with lung.ab,ti.
- 22 carcinoma with lung.ab,ti.
- 23 16 or 17
- 24 18 or 19 or 20 or 21 or 22
- 25 exp Drug Therapy/
- 26 drug therapy.ab,ti.
- 27 chemotherapy.ab,ti.
- 28 25 or 26 or 27
- 29 exp radiotherapy/
- 30 radiotherapy.ab,ti.
- 31 29 or 30
- 32 exp surgery/
- 33 surgery.ab,ti.
- 34 32 or 33
- 35 28 or 31 or 34
- 36 6 or 15
- 37 36 and 24 and 35

Appendix C Suggested coding

Please provide data on all patients randomised. You may complete data forms (provided on request) or supply your data as a computer printout, on floppy disk (formatted for PC) or by email. Data can be in almost any format (ASCII, Excel, Dbase, FoxPro etc.), but please indicate which format has been used. It would be helpful if you used the coding suggested, however you may code the data in the way that is most convenient for you. Please supply us with full details of the data coding system used. If sending data via email, please encrypt the data and let us know how it has been encrypted in a separate email.

Patient Identifier	D. C. I.I.			
	Preferably not na Type Width	i me character 15		
Date of Birth				
	Type Width Code	date 8 or 6 date in dd/mm/yy	yy (recommended)	or dd/mm/yy format
Age	Type Width	numeric 3		
	Code	age in years unknown = 999		
Sex				
	Type Width Code	numeric 1 1=male 2=female 9=unknown		
Tumour stage used				
J	Type Width Code	numeric 1 1=pTNM 2=AJCC 3=1986 ISS 4=1997 UICC		
If pTNM used Tumour Stage pTNM				
Tumour Stage privin	Type Width	character 3	aNI Stano	
	Code	pT Stage 0=pT0 X=pTX S=pTis 1=pT1 2=pT2 3=pT3 4=pT4 9=Unknown	pN Stage 0=pN0 1=pN1 2=pN2 3=pN3 9=Unknown	pM stage 0=pM0 1=pM1 9=Unknown
If AJCC used Tumour Stage AJCC				
Tumour Staye ASCC	Type Width	numeric 1		

	Code	1=stage 1 2=stage 2 3=stage 3 4=metastatic 9=unknown
If ISS used		
Tumour Stage 1986 ISS	Type Width Code	numeric 1 1=stage 1 2=stage 2 3=stage 3A 4=stage 3B 5=stage 4 9=unknown
If 1997 staging used		
Tumour Stage 1997 UICC	Туре	numeric
	Width	1
	Code	1=stage 1A 2=stage 1B 3=stage 2A 4=stage 2B 5=stage 3A 6=stage 3B 7=stage 4 9=unknown
Histology		
	Type Width Code	numeric 1 1=small cell 2=adenocarcinoma 3=squamous 4=mixed 5=large cell undifferentiated 6=NSC unspecified 7=other 9=unknown
Performance Status (Karno	fsky)	
	Type Width Code	numeric 3 10-100 999=unknown
Performance Status (WHO/	ECOG)	
	Type Width Code	numeric 1 1-4 9=unknown
Treatment Allocated		
	Type Width Code	numeric 1 1=supportive care 1=supportive care + chemotherapy If more than 1 regimen of chemotherapy used, please specify and use 3,4,5 etc for each different regimen

Date of Randomisation	Type Width Code	date 8 or 6 date in dd/mm/yyyy (recommended) or dd/mm/yy format
Survival Status	Type Width Code	numeric 1 0=alive 1=dead
Date of Death / Last Follow-up	Type Width Code	date 8 or 6 date in dd/mm/yyyy (recommended) or dd/mm/yy format
Cause of Death	Type Width Code	numeric 1 1=lung cancer 2=treatment related 3=other 9=unknown
Excluded	Type Width Code	numeric 1 0=included in analysis 1=excluded from analysis 9=unknown
Reason for Exclusion	Type Width	character 25

Appendix D

Completed Individual Patient Data Meta-analyses by MRC and IGR

Meta-analysis Group, MRC Clinical Trials Unit, UK

Advanced Ovarian Cancer Trialists Group. Chemotherapy in advanced ovarian cancer: an overview of randomised clinical trials. British Medical Journal 1991;303:884-93.

Advanced Bladder Cancer Overview Collaboration. Does neo-adjuvant cisplatin-based chemotherapy improve the survival of patients with locally advanced bladder cancer: a metaanalysis of individual patient data from randomised clinical trials. British Journal of Urology 1995;75:206-13.

Non-small Cell Lung Cancer Collaborative Group. Chemotherapy in non-small cell lung cancer: a meta-analysis using updated data on individual patients from 52 randomised clinical trials. British Medical Journal 1995;311:899-909.

Sarcoma Meta-analysis Collaboration. Adjuvant chemotherapy for localised resectable soft tissue sarcoma in adults: meta-analysis of individual patient data. Lancet 1997;350:1647-54.

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PORT Meta-analysis Trialists Group. Postoperative radiotherapy in non-small-cell lung cancer: systematic review and meta-analysis of individual patient data from nine randomised controlled trials. The Lancet 1998;352:257-63.

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Advanced Bladder Cancer (ABC) Meta-analysis Collaboration. Neoadjuvant chemotherapy in invasive bladder cancer: a systematic review and meta-analysis. Lancet 2003;361:1927-34.

Neoadjuvant Chemotherapy for Cervical Cancer Meta-analysis Collaboration. Neoadjuvant chemotherapy for locally advanced cervical cancer: a systematic review and meta-analysis of individual patient data from 21 randomised trials. European Journal of Cancer 2003;39:2470-86.

Service de Biostatistique et d'Epidemiologie, Institut Gustave-Roussy, France

Pignon J-P, Arriagada R, Ihde DC, Johnson DH, Perry MC, Souhami RL, et al. A meta-analysis of thoracic radiotherapy for small-cell lung cancer. New England Journal of Medicine 1992;327:1618-24.

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Pignon JP, Bourhis J, Domenge C, Designé L, on behalf of the MACH-NC Collaborative Group. Chemotherapy added to locoregional treatment for head and neck squamous cell carcinoma: three meta-analyses of updated individual data. Lancet 2000;355:949-55.

Bourhis J, Syz N, Overgaard J, Ang KK, Dische S, Horiot J, et al. Conventional vs modified fractionated radiotherapy. Meta-analysis of radiotherapy in head & neck carcinoma: a meta-analysis based on individual patient data. International Journal of Radiation Oncology Biology Physics 2002;54(Suppl):71-2.

Piedbois P, Michiels S for the Meta-analysis Group in Cancer. Survival benefit of 5FU/LV over 5FU bolus in patients with advanced colorectal cancer: an updated meta-analysis based on 2751 patients. Proceedings of the American Society of Clinical Oncology 2003;22:294.

Auperin A, Le Pechoux C on behalf of the MAC3-LG Group. Meta-analysis of randomized trials evaluating cisplatin or carboplatin-based concomitant chemoradiation versus radiotherapy alone in locally advanced non-small cell lung cancer (NSCLC). Lung Cancer 2003;41(Suppl 2):S69.