

AN UNUSUAL STALK ROT OF SUGARCANE CAUSED BY *PHAEOCYTOSTROMA SACCHARI* IN THE KWAZULU-NATAL MIDLANDS

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Abstract

A severe and unusual stalk rot was first observed in the KwaZulu-Natal Midlands in October 1998 after prolonged dry conditions during winter and spring. The rot was subsequently found to be common in many fields of a number of varieties. In a survey in the area, 64 out of 137 fields were found to be affected to some degree. An orange-brown rot occurred in more than 50% of the internodes in most of the affected stalks. The rot occurred in both young and old cane of varieties N12, N16 and N21. The purity of infected stalks was reduced, and this caused a number of cane consignments to be rejected by the mills. The dominant fungal pathogen isolated from the affected stalks was identified as *Phaeocytostroma sacchari* (Ellis and Everh.) Sutton. This pathogen also causes rind disease, a common but minor infection of the rind of over-mature and moribund stalks, but one that has been reported to progress to cause severe stalk rotting, if mature cane is subjected to prolonged poor growing conditions such as drought. This is the first record of rind disease progressing to cause a severe stalk rot in the South African sugar industry.

Keywords: Sugarcane, stalk rot, *Phaeocytostroma sacchari*, rind disease

Introduction

Phaeocytostroma sacchari (Ellis and Everh.) Sutton causes rind disease, which is a common but usually minor condition that affects cane stalks weakened by insect injury or other wounds, cane growing under unfavourable conditions such as drought, and over-mature and moribund stalks (Johnston, 1917; Abbott *et al.*, 1964). The diagnostic symptom of rind disease is the pustules, which appear as coiled, black masses of spores under humid conditions. The pustules break through the surface of the rind of affected stalks, and can also be present on leaf sheaths and midribs (Abbott *et al.*, 1964).

When there is severe drought, particularly late in the growing season, rind disease may develop into a stalk rot, which can cause substantial sugar losses, particularly in susceptible varieties (Liu *et al.*, 1977). Under these adverse conditions an internal rotting appears in the mature portions of growing stalks, followed by a discolouration of the rind and develop-

ment of pustules under appropriate conditions. If the drought persists, the portion of the stalk above the infected internodes may continue to deteriorate and become desiccated. In severe infections, rotting can extend into the stubble and kill the entire stool (Abbott *et al.*, 1964). The stalk rot phase of the disease is uncommon and had not previously been reported from South Africa.

In October 1998, an unusual and severe stalk rot of unknown identity was reported in a field of 18 month old cane of variety N12 in the Midlands South extension area. This rot was subsequently found to be common in other fields in the area. A number of infected cane consignments were rejected by the mills because of low purity.

The aims of this study were to conduct disease surveys in the Midlands to estimate the extent of the unusual stalk rot and to identify the causal organism.

Materials and Methods

Field survey

A survey was conducted in the Midlands South extension area, where the disease was first reported. A total of 137 fields on 57 farms were sampled in October and November 1998. Most of the surveying was conducted by the field inspection team of the Local Pest and Disease Control Committee (LP&DCC). Samples consisting of 100 randomly selected stalks were taken from each field. The stalks were split and examined for borer damage and rotting. Stalk rotting was identified as being due either to red rot (*Glomerella tucumanensis*), based on its characteristic symptoms, or 'new' stalk rot. The latter was usually accompanied by a distinctive sour odour.

Further stalk samples were taken from 15 affected fields and examined in the Pathology laboratory at the SASA Experiment Station (SASEX). These stalks were examined for external damage and symptoms of red rot and the new stalk rot. The number of internodes affected by the stalk rot was also recorded.

Rainfall

Rainfall figures were supplied by SASEX from data collected from meteorological stations in the Midlands area.

Pathogen isolation

Between two and five stalks were taken from each of the samples that were brought to SASEX. The stalks were split in half longitudinally to assess the extent of rotting and 1 cm³ pieces of internal internodal tissue were cut from the interface between infected and apparently healthy tissue. The pieces were surface sterilised by soaking in 0,35% sodium hypochlorite (10% 'Jik') for five minutes and the outer surfaces were aseptically removed. The remaining tissue pieces (approximately 0,5 cm³) were again surface sterilised in sodium hypochlorite for 2 minutes, air dried on sterile tissue paper and dipped in 70% ethanol for 2 minutes before flaming and plating onto potato dextrose agar (PDA). Mycelium growing from the tissue sections was immediately transferred onto fresh PDA. All petri dishes were kept on a laboratory bench at 20-25°C under natural light. Most of the cultures appeared to be *Phaeocystostroma sacchari*, and some a *Fusarium* sp.

Pathogenicity tests

Stalks of sugarcane variety N12 were collected from the field and cut into approximately 10 cm long segments that were quartered longitudinally. The pieces were placed into test tubes and autoclaved twice at 121°C for 20 minutes. Each sterile stalk piece was then inoculated with a disc of agar cut from each culture. The tubes were left on a laboratory bench and examined periodically.

The marcotting procedure used by the SASEX Plant Breeding Department (Anon., 1975; Nuss, 1977) was used to obtain single, live sugarcane stalks that could be inoculated with fungal cultures and grown in the glasshouse. Stalks of varieties N12 and NCo376 were cut just above the soil surface in the field. The basal part of each stalk was placed in and through a metal cylinder (40 cm long, 15 cm diam.) that was then filled with a growing medium of composted bagasse and filtercake. Approximately 15 cm of the stalk extended below the cylinder and this was placed in a trough containing a sulphur dioxide solution. Once roots had developed within the cylinder, the trough was removed and the stalk was watered through the growing medium.

A 1,5 mm drill bit was used to make a horizontal hole in each marcotted stalk, midway between the nodes of the first full internode above the growing medium. The hole was drilled horizontally to the centre of the stalk tissue. Fruiting bodies (sclerotia) from ten cultures of *P. sacchari* and spores from one culture of the *Fusarium* sp were suspended in 5 ml sterile distilled water in McCartney bottles. The bottles were vortexed to release spores from the sclerotia. Each plant was inoculated with 0,5 ml of spore suspension using a 1 ml hypodermic needle. The holes were then sealed with parafilm.

This experiment consisted of 12 treatments, each replicated four times for each variety. A total of 80 stalks were inoculated with *P. sacchari*. Control stalks were injected with sterile distilled water.

After inoculation the plants were watered daily for five

weeks and then water was withheld for two weeks. After the seven week period external symptoms were noted and the stalks were cut in half longitudinally and the number of infected internodes counted. Pathogens that has colonised the stalk tissues of variety N12 were identified as described above under 'Pathogen isolation'.

Results and Discussion

Field survey

Sixty-four (47%) of the 137 fields sampled were infected with stalk rot to some degree (Table 1). Forty-one of the 64 affected fields had less than 10% infected stalks. Seven fields contained between 25 and 50% affected stalks, in nine fields more than 50% of the stalks were infected, and in one field all the stalks sampled were infected. In most cases, not all the stalks in one stool were infected.

Table 1. Number of fields and percent stalks affected by stalk rot in the Midlands South extension area.

% stalks with rotting	No. of fields
0	73
1-5	30
6-10	11
11-25	7
26-50	7
51-75	6
76-100	3
Total fields	137

The internal tissues of affected stalks were red/orange in colour (Figure 1) and had a distinctive sour odour. The colour was easily distinguished from the brighter red rotting caused by red rot. In most stalks, cushiony masses of mycelium were present, as described by Martin (1938). Hair-like masses of spores were visible emerging from pustules on the rind and leaf sheaths of some stalks (Figure 2). The purity of infected stalks was reduced, causing a number of cane consignments to be rejected by the mills.

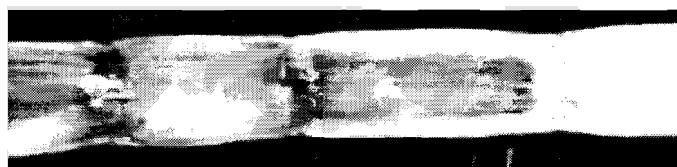


Figure 1. Internal stalk rotting of N12 caused by *Phaeocystostroma sacchari*.

Variety N12 is extensively grown in the Midlands South area and therefore it constituted a large proportion of the fields sampled (Table 2). The stalk rot was not limited to N12, and varieties N16, N21 and NCo376 also had symptoms. Of the samples examined at SASEX, a mean of 73% of the internodes in affected stalks were damaged (Table 3).

Infection was not associated with borer or other damage. In most stalks, infection appeared to have entered the stalk in the region of the third or fourth internode, possibly through the leaf scars, buds or root primordia.

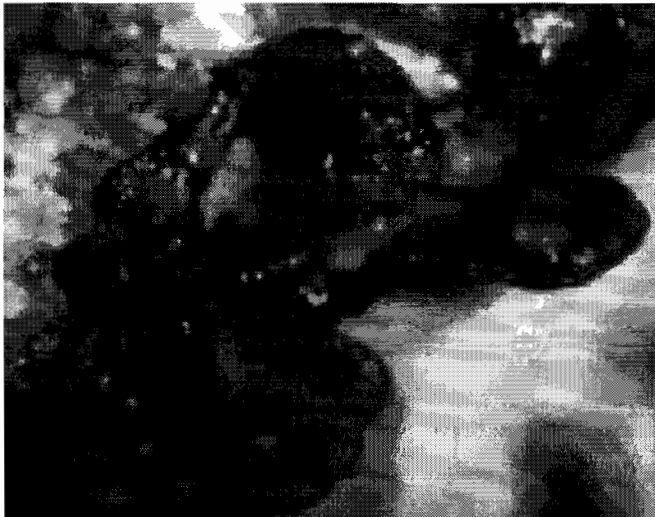


Figure 2. Hair-like spore masses emerging from rotted stalks.

Table 2. Incidence of stalk rot in different varieties grown in the Midlands South extension area.

Variety	No. fields sampled	No. (and %) fields with stalk rot
N11	1	0 (0)
N12	111	55 (50)
N16	17	5 (29)
N21	3	2 (67)
NCo376	5	2 (40)
Total fields	137	-

Table 3. Percent internodes damaged in stalks found to be infected with stalk rot.

% internodes affected	No. of samples
1-25	0
26-50	1
51-75	6
76-100	8
Total samples	15

By February 1999, all affected fields had been harvested. A 600 stalk sample taken at that time from a field of N12 showed that the stalk rot was still present in the area but at low levels, with only 1,6% of the stalks sampled being affected. Further surveys of previously affected fields are planned for the coming season to determine whether the disease is still prevalent.

Rainfall data

The Midlands area received below average rainfall from June to November 1998 (Figure 3). September was particularly dry, with only 24 mm (39% LTM) being recorded at three meteorological stations in the Midlands South area and 32 mm (46% LTM) recorded at six sites in the Midlands North area. Prolonged periods of drought are reported to have caused severe outbreaks of stalk rot caused by *P. sacchari* in Queensland (Abbott *et al.*, 1964) and Hawaii (Anon, 1958). The prolonged and unusually intense drought that occurred in the Midlands region of the South African sugar industry in mid-1998 is therefore thought to be the main rea-

son that *P. sacchari*, usually regarded as a common but minor pathogen, caused severe damage to cane in 1998.

Pathogen isolation and identification

Sixty-four fungal cultures were isolated, seven of which were identified as a *Fusarium* spp. and 57 as *P. sacchari*, as confirmed by one of the authors (CR) at the National Collection of Fungi Biosystemics Division. *Fusarium* sp. and *P. sacchari* occurred together on four tissue segments. In pure culture on PDA, the mycelium of *P. sacchari* was dark grey. Black, spherical fruiting bodies (conidiomata) 1-2 mm in diameter, appeared in culture after seven days. When the conidiomata were viewed under the light microscope a number of pycnidia were seen in each. Each pycnidium contained masses of one-celled, light brown, cylindrical spores measuring 10-13,8 µm x 3,8 µm (Figure 4). Square crystals of calcium oxalate, measuring between 10 and 12,5 µm, were also observed.

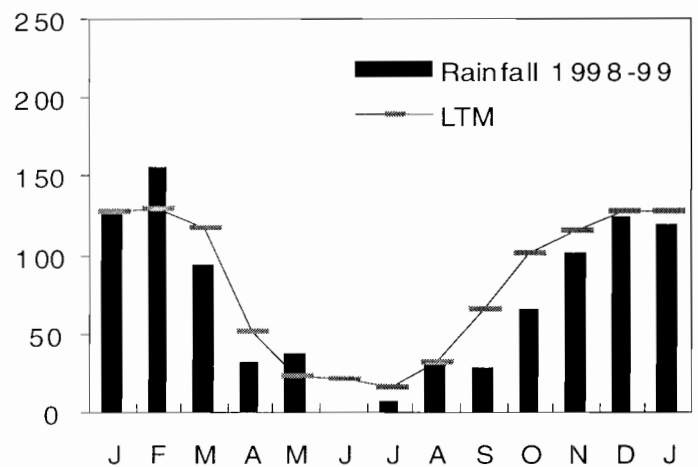


Figure 3. Mean monthly rainfall and long term mean rainfall (LTM) in the Midlands region of KwaZulu-Natal, January 1998 - January 1999 (mm).

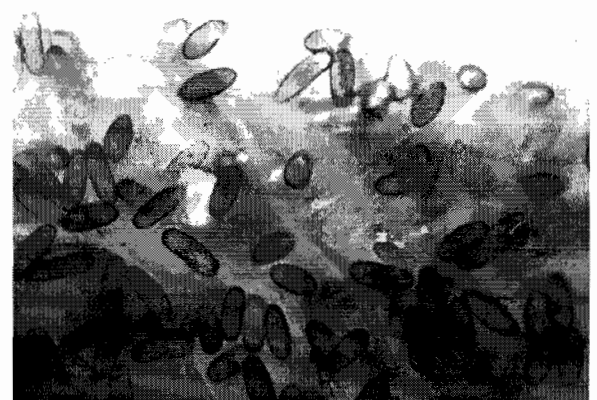


Figure 4. Spores of *P. sacchari*.

Pathogenicity tests

In the laboratory, following inoculation and incubation at room temperature, all stalk sections inoculated with *P. sacchari* were soon enveloped by mycelium. Initially, white spots developed on the rind. A few weeks later black spore masses emerged from these areas (Figure 5). Some of the spore masses were coiled, as described by Abbott *et al.* (1964). This confirmed that the predominant fungus isolated from the rotted portions of the stalks was *P. sacchari*, the causal organism of rind disease. The stalk pieces inoculated with *Fusarium* were covered with a dark purple mycelium.

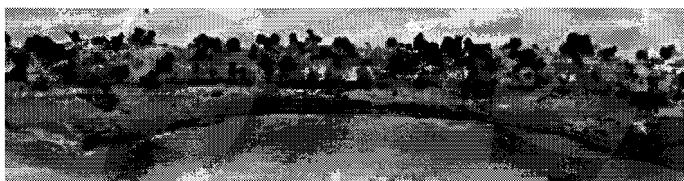


Figure 5. Black spore masses of spores of *P. sacchari* emerging from pustules on rotted stalks after inoculation.

In the glasshouse experiment, when most stalks inoculated with *P. sacchari* and the untreated controls were split, only the inoculated internodes were reddened. This discolouration spread to the upper internodes in only nine of the inoculated stalks (six stalks had two infected internodes, two stalks had three infected internodes and one stalk had six infected internodes). This result showed that the inoculation technique was unsuccessful in most stalks. This may have been due to a number of factors, including the possibility that the inoculum density of the fungal spore suspension may have been too low. It is also probable the stalks were not stressed long enough after inoculation for the infection to spread throughout the stalk. Further experiments may be necessary to determine the length of drying-off period for symptoms to occur.

It may be necessary to evaluate other methods for pathogenicity tests for *P. sacchari*. For example, Liu *et al.* (1977) successfully inoculated four-budded setts by spraying spores onto the stalk surface before covering with a plastic bag for two to three days; after ten days 93% of the varieties were rated as having an intermediate or susceptible reaction to the fungus.

Twenty days after the reddened, inoculated internodes or uppermost infected internodes of N12 from the glasshouse experiment were quartered, surface sterilised and placed into test tubes, most of the 40 sections that had been inoculated with *P. sacchari* showed symptoms of rind disease. This result confirms that the fungus was present in the actively

growing marcotted stalks, although conditions in the glasshouse were not conducive to the growth and spread of the fungus within the stalks.

Conclusions

A severe stalk rot occurred in many sugarcane fields in the KwaZulu-Natal Midlands in late 1998, after prolonged dry conditions during the preceding winter and spring months, and caused substantial economic damage. The stalk rot was caused by the fungal pathogen *Phaeocystostroma sacchari*. This pathogen also causes rind disease of sugarcane, a common but usually minor infection of the rind of over-mature or moribund stalks. The outbreak of severe stalk rot in the Midlands was the first record of this phase of the disease in South Africa, and confirms the few published reports that *P. sacchari* can cause severe damage to sugarcane crops that are subjected to prolonged poor growing conditions, such as drought. This new outbreak was unusual in that it was common, it affected a number of varieties, and it affected crops with a wide range of ages and was not confined to over-mature cane.

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THE INCIDENCE AND EFFECTS OF RATOON STUNTING DISEASE OF SUGARCANE IN SOUTHERN AND CENTRAL AFRICA

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Abstract

Surveys have shown that ratoon stunting disease (RSD) is common in most sugar industries in southern and central Africa. In South Africa in 1998, approximately 12% of commercial cane fields contained some level of RSD infection, but the mean number of stalks infected was low. The most recent estimates of the numbers of infected fields in other industries are: Swaziland 30%; Zambia 50%; Kenya, Malawi, Uganda and Zimbabwe 60-90%; Tanzania and Mafambisse estate in Mozambique 100%. In some of these industries the majority of stalks within fields are infected. Field experiments have shown that RSD can cause reductions in yield of 15-30% under good irrigated conditions, and 20-40% under average rainfed growing conditions in varieties that are widely grown in Africa. By integrating survey data and experiment results, it is estimated that yield losses due to RSD in South Africa are equivalent to approximately 1% of current production, but that losses of 10-20% or even greater are probable in some of the other industries.

Minimising the effects of RSD on production should be a priority throughout the region. To achieve this, attention must be given to two main factors. Firstly, the consistent production of healthy seedcane through well-managed schemes incorporating hot water treatment and, secondly, improving the efficiency of stubble destruction before fields are replanted, to prevent RSD surviving in infected volunteer plants and spreading to new plantings. In many areas this will require longer breaks from cane before fields are replanted.

Key words: ratoon stunting disease, RSD, hot water treatment, yield loss, *Clavibacter*

Introduction

The South African Sugar Association Experiment Station (SASEX) has conducted large scale surveys of RSD incidence in the South African sugar industry since 1977. A number of other sugar industries in Africa have made use of the SASEX diagnostic service. Survey data from the mid to late 1990s are available from Swaziland, Malawi, Zambia, Kenya, Tanzania, Uganda and Mozambique. Information on the status of RSD in Zimbabwe from 1996 to 1998 is available from surveys by the Zimbabwe Sugar Association Experiment Station (ZSAES). From the time of the earliest surveys it was apparent that high levels of RSD occurred in

certain parts of the South African sugar industry and in most of the other industries.

In most sugarcane industries in Africa production is based mainly on SASEX-bred varieties. The effects of RSD on these varieties are routinely determined by SASEX in field experiments. The magnitude of losses in many varieties when infected by RSD is known to be large under both rainfed and irrigated conditions.

An important feature of sugarcane production in most African sugar industries is that, traditionally, only short breaks from cane, sometimes as little as two to four weeks, are applied between destroying old crops and replanting fields. Consequently, numerous volunteer plants survive from old crops into new plantings.

This paper summarises the status of RSD in industries where surveys have been conducted. By considering survey data together with estimates of yield loss in controlled experiments, estimates of the effect of RSD on production in different countries are possible. These estimates are presented and key factors necessary to reduce the economic effects of RSD are identified.

Survey methods

Surveys of RSD incidence in South Africa have been conducted by SASEX since 1977. Currently, samples from approximately 7 000 fields are tested annually. These include the majority of seedcane sources intended for planting. Additionally, in all mill supply areas large numbers of commercial fields are selected randomly for survey purposes.

Until recently, surveys of RSD in South Africa were based on the examination, by phase contrast microscopy (PCM), of xylem sap extracted from stalks collected in the field (Bailey and Fox, 1984). PCM is also used on a large scale in Zimbabwe. In early 1998, routine diagnosis in South Africa was changed to an evaporative binding-enzyme immunoassay (EB-EIA), based on a method developed in Australia (Croft *et al.*, 1994) and using a polyclonal antiserum to a local isolate of *Clavibacter xyli* subsp. *xyli* (Cxx).

RSD diagnosis in Zimbabwe in the period 1996-98 was based on PCM (Zvoutete, personal communication¹).

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Samples from Swaziland and Mozambique were examined by either PCM or immunofluorescence microscopy (IFM). Material from all the other countries was examined by IFM, again using a polyclonal antiserum to C.x.x. For IFM, drops of xylem sap extracted from stalks are dried onto multi-celled slides. These are then delivered to SASEX for processing and examination.

In most cases, samples consisted of 20 stalks per field or section of field, chosen from weaker plants. This biased sampling was intended to increase the likelihood of detecting RSD, if present in the field. Larger, randomly collected samples of up to 100 stalks were used to determine the percentage of infected stalks in fields in South Africa. Estimates of the percentage of infected stalks per field were possible for several other countries, including Zimbabwe.

Effect of RSD on yield

Numerous field experiments have been conducted by SASEX to determine the effects of RSD on cane and sugar yields. In the widely grown variety NCo376, which is the main variety in a number of African sugar industries, losses in sugar yield of 20-40% under average rainfed conditions and up to 20% under good irrigated conditions have been recorded. Some varieties, such as N17 and N14, are more intolerant than NCo376 and substantial losses have been recorded in most of the large number of varieties that have been tested (Bailey and Bechet, 1986; 1995). The results of two recent trials are shown in Figure 1.

Rainfall in southern and central Africa is mainly seasonal and often erratic. When infected crops are subjected to moisture stress from drought or inadequate irrigation, losses due to RSD can be dramatic. Reductions in sugar yield of 76% were recorded in variety NCo376 in the drought season of 1980-81 (Bailey and Bechet, 1986).

Incidence of RSD and effects on production

South Africa

Surveys in the late 1970s showed that approximately 30% of fields in the southern, rainfed part of the South African industry contained some level of RSD. The incidence was greater in the northern production area, where the crop is grown under full irrigation, similar to conditions in most of the other industries discussed. In this northern area in the early 1980s, 62% of fields in Mpumalanga and 40-50% of fields in the Pongola mill area were infected (Bailey and Fox, 1984; Bailey and Tough, 1991).

In the last two decades in most parts of the South African industry, the mean number of fields in which RSD was detected has declined steadily. In 1997, RSD was detected in 21% of fields in the northern area and 6% of fields in the southern area, and the industry mean was 9%. This was the lowest ever recorded. The situation in the industry as a whole in 1997 is illustrated in Figure 2. Using EB-EIA, the mean number of fields infected in 1998 was estimated to be 12%.

In the southern, rainfed area (approximately 78% of total South African sugar production), intensive surveys have shown that the mean number of infected stalks in fields where RSD is present is low, approximately 1%. With this mean level of stalk infection, yield losses in most parts of the southern area are now thought to be negligible. The estimated mean number of stalks infected in the northern production area in 1997 was 7%, and the industry mean was 2%. In the South African industry as a whole, losses in 1979 were estimated to be approximately 5% of annual production (Bailey, 1979), but current losses are estimated to be equivalent to approximately 1% of production.

Swaziland and Zimbabwe

A number of RSD surveys based on IFM (1992-96) and

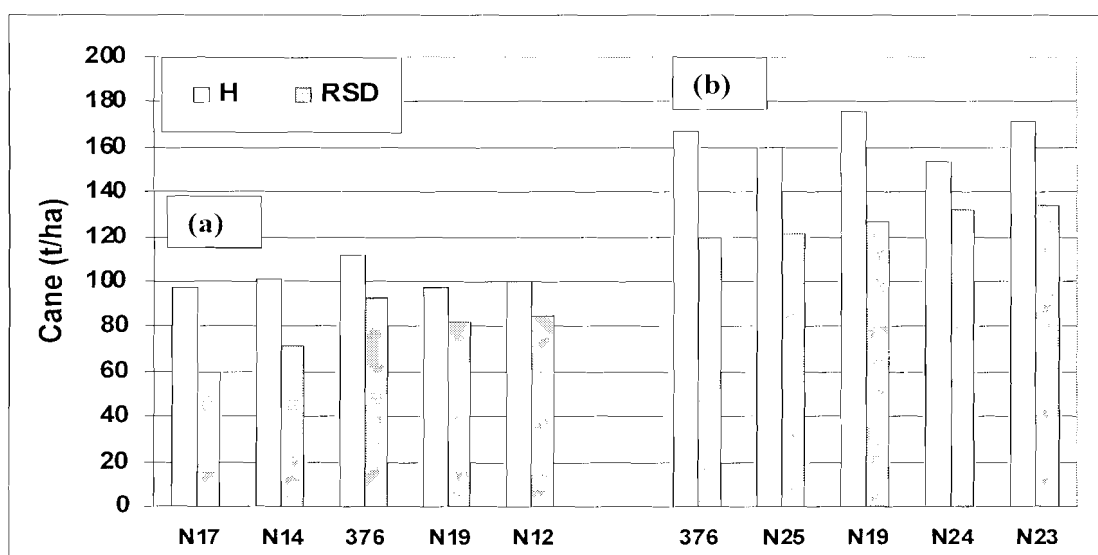


Figure 1. Effect of RSD on cane yield under (a) rainfed conditions at Mount Edgecombe (2R, 13.5 months, 1996); (b) irrigated conditions at Pongola (2R, 11.7 months, 1997; H = healthy seedcane, RSD = infected seedcane).

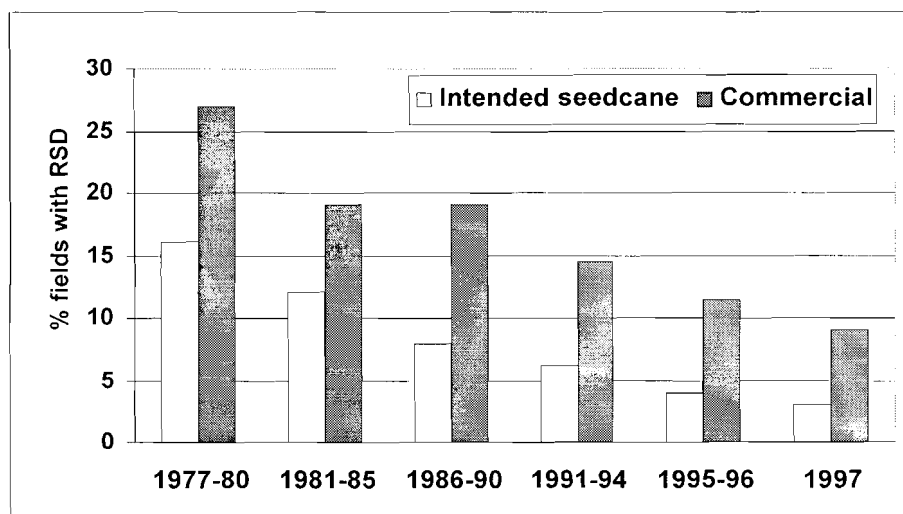


Figure 2. Mean incidence of RSD in intended seedcane sources and commercial fields in the South African sugar industry, 1977-1997 (% fields in which RSD was detected); any intended seedcane found to have RSD is not planted.

Table 1. RSD incidence in Swaziland, 1992-1998 (% fields with RSD).

Year	No. fields tested	% fields with RSD
1992	159	22
1993	291	15
1995	315	15
1996	247	30
1998	330	28

PCM (1998) have been conducted in Swaziland by SASEX. The mean number of commercial fields in which RSD was recorded ranged from 15 to 30% (Table 1), partly depending on the areas surveyed, but the two latest surveys gave similar results – means of approximately 30% infected fields (Table 1). Losses in production due to RSD are estimated to be approximately 4% of current production.

Until recently there had been little use of hot water treatment (HWT) in Zimbabwe for many years. After a devastating drought in the early 1990s the entire industry was re-established, starting in 1993. Unfortunately, the scarce seedcane stocks that were available had high levels of RSD and the replanting exercise served to spread RSD throughout the industry. Surveys of RSD incidence have been conducted by ZSAES since 1995 using PCM (Zvoutete, personal communication). Survey data from 1998 show that more than 80% of commercial cane fields were infected with RSD, and that approximately 60% of stalks were infected (Table 2). Reductions in yield in Zimbabwe due to RSD are estimated

to be approximately 10% of annual production. The situation with regard to seedcane offers some encouragement, with 20% of intended sources being infected.

Other countries

Surveys in other countries have been less frequent, but it is likely that the data recorded are representative of the current situations. Extremely high levels of RSD were recorded at both Dwangwa and Nchalo in Malawi, at Nakambala in Zambia, at both Mumias and South Nyanza in Kenya, and at Kinyara in Uganda (Table 3). The results indicated that the majority of stalks in fields where RSD was identified were infected. In all these industries, the use of HWT to eliminate RSD from seedcane stocks has been sporadic. A feature of cane production in most of these industries is the relatively short break from cane before fields are replanted. Losses in yields in these industries probably amount to 10-20% of production.

Samples from 24 fields at Kilombero in Tanzania were

Table 2. RSD incidence in Zimbabwe, 1996-98 (% fields infected and mean % stalks infected for all fields).

Year	No. fields tested	% fields with RSD	% stalks infected (all fields)
1996	612	92	71
1997	1 372	64 ¹	51 ¹
1998	867	82	58

Note: ¹ the apparent lower incidence in 1997 was due to many samples being from seedcane and plant cane fields.

Table 3. Incidence of RSD in Zambia, Malawi, Kenya and Uganda, 1993-1998 (% fields with RSD).

Country, area & year	No. fields tested	% fields with RSD
Zambia (Nakambala,)		
1994	196	98
1996 + 1997	161	54
1998	50	46
Malawi		
Dwangwa (1993)	36	67
Nchalo (1995)	62	74
Kenya		
South Nyanza (1993)	196	96
Mumias (1993)	391	57
Uganda		
Kinyara (1997)	50	82
Tanzania		
Kilombero (1998)	24	100
Mozambique		
Mafambisse (1998)	21	100

examined in 1998. All were found to be infected and the results indicated that most stalks were infected. Information on the status of RSD at Mafambisse in Mozambique was obtained by one of the authors (RAB) on a visit in 1991. Prior to that there had been no HWT on the estate for many years. On inspecting fields, the internal nodal symptoms of RSD were invariably found. Samples from a number of fields were brought to SASEX for confirmation by PCM. All were found to be infected and it was concluded that RSD was ubiquitous on the estate. These results were confirmed by further sampling in 1998. If the survey results are representative, it is estimated that reductions in production at both Kilombero and Mafambisse exceed 20%. In contrast, no RSD was found in stocks of N19 on a visit to Maragra estate in southern Mozambique in 1997. This estate is now being rehabilitated.

Discussion and conclusions

Except for South Africa, RSD is estimated to be having a significant impact on sugar production in all the countries in southern and central Africa from which survey data are available.

Good progress has been made in reducing the incidence and economic significance of RSD in South Africa. The main factor that has contributed to the improved situation in all parts of the industry has been the widespread use of seedcane production schemes based on HWT. Most of the seedcane now planted is obtained from sources that have been tested for freedom from RSD. In all areas, there has been a close correlation between the RSD status in commercial fields and the quality of the seedcane planted. Further contributing factors include the improved attention given by growers to stubble destruction and the greater use of longer breaks from cane between plantings. It is recommended that fields have a break from cane of at least three months before replanting.

An important factor that contributed to RSD control in South Africa was the widespread publicity given to research results on the effect of RSD on yields and on the rate at which RSD

can spread (Bailey and Tough, 1992). The diagnostic service has provided valuable information on the RSD situation at farm, area and industrial levels, as well as providing specific information to aid growers in making practical management decisions concerning seedcane sources and plough-out fields.

Sugarcane in Swaziland is produced mainly on large miller-cum-planter estates. As in most other industries in the region, the period between stubble destruction and replanting is usually short, sometimes only a few weeks. The industry has had a national seedcane scheme since the late 1970s, operated by the Swaziland Sugar Association. In this, elite seedcane is produced after HWT in an area remote from the main production areas and is used to establish commercial nurseries on the estates. This has been the main factor in achieving substantial control of RSD compared with most other industries in the region. Further progress in Swaziland below the current level of approximately 30% fields infected requires greater attention to stubble destruction. This will necessitate longer breaks between plantings than is currently practised.

In Zimbabwe, control of RSD has been identified as a high priority within the last three years, and a major drive to improve seedcane quality by greater use of HWT is now in progress on most estates. The diagnostic service provided by ZSAES will be a key factor in achieving success.

Major factors impeding an improvement in the RSD situations in the other industries mentioned are the current lack of HWT facilities and the lack of local services for large scale diagnosis. SASEX can provide further assistance with diagnosis but the development of local expertise would assist the situation.

In all the industries where RSD occurs at high levels, attention must be given to increasing the duration of the break between plantings to minimise the survival of infected volunteer regrowth. This recommendation is likely to meet some resistance until its value is supported by local survey and research results.

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